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(54) POLYNUCLEOTIDE ENCODING A POLYPEPTIDE HAVING HEPARANASE ACTIVITY AND **EXPRESSION OF SAME IN TRANSDUCED CELLS** 

POLYNUKLEOTID KODIEREND FÜR EIN POLYPEPTID MIT HEPARANASE-AKTIVITÄT UND DESSEN EXPRESSION IN TRANSDUZIERTEN ZELLEN

POLYNUCLEOTIDE CODANT POUR UN POLYPEPTIDE PRESENTANT UNE ACTIVITE D'HEPARANASE ET SON EXPRESSION PAR DES CELLULES TRANSDUCTEES

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(73) Proprietors:

· Insight Biopharmaceuticals Ltd 76121 Rehovot (IL)

 HADASIT MEDICAL RESEARCH SERVICES AND DEVELOPMENT LTD. 91120 Jerusalem (IL)

(72) Inventors:

 PECKER, Iris 75204 Rishon Le Zion (IL)

 VLODAVSKY, Israel 90805 Mevaseret Zion (IL)

 FEINSTEIN, Elena 76214 Rehovot (IL) (74) Representative: Wachenfeld, Joachim **Vossius & Partner** Postfach 86 07 67 81634 München (DE)

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 DATABASE GENBANK ON STN, US NATIONAL LIBRARY OF MEDICINE, (Bethesda, MD, USA), No. N32056, HILLIER et al., "The WashU-Merck EST Project", 10 January 1996, XP002915282

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- DATABASE GENBANK ON STN, US NATIONAL LIBRARY OF MEDICINE, (Bethesda, MD, USA), No. N30845, HILLIER et al., "The WashU- Merck EST Project", 05 January 1996, XP002915283
- DATABASE GENBANK ON STN, US NATIONAL LIBRARY OF MEDICINE, (Bethesda, MD, USA), No. N30824, HILLIER et al., "The WashU-Merck EST Project", 05 January 1996, XP002915284
- DATABASE GENBANK ON STN, US NATIONAL LIBRARY OF MEDICINE, (Bethesda, MD, USA), No. AA304653, ADAMS et al., "Initial Assessment of Human Gene Diversity and Expression Patterns Based Upon 83 Million Nucleotides of cDNA Sequence", 18 April 1997, XP002915285

## Description

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## FIELD AND BACKGROUND OF THE INVENTION

[0001] The present invention relates to a polynucleotide, referred to hereinbelow as *hpa*, encoding a polypeptide having heparanase activity, vectors including same and transduced cells expressing heparanase. The invention further relates to a recombinant protein having heparanase activity.

[0002] Heparan sulfate proteoglycans: Heparan sulfate proteoglycans (HSPG) are ubiquitous macromolecules associated with the cell surface and extra cellular matrix (ECM) of a wide range of cells of vertebrate and invertebrate tissues (1-4). The basic HSPG structure includes a protein core to which several linear heparan sulfate chains are covalently attached. These polysaccharide chains are typically composed of repeating hexuronic and D-glucosamine disaccharide units that are substituted to a varying extent with N- and O-linked sulfate moieties and N-linked acetyl groups (1-4). Studies on the involvement of ECM molecules in cell attachment, growth and differentiation revealed a central role of HSPG in embryonic morphogenesis, angiogenesis, neurite outgrowth and tissue repair (1-5). HSPG are prominent components of blood vessels (3). In large blood vessels they are concentrated mostly in the intima and inner media, whereas in capillaries they are found mainly in the subendothelial basement membrane where they support proliferating and migrating endothelial cells and stabilize the structure of the capillary wall. The ability of HSPG to interact with ECM macromolecules such as collagen, laminin and fibronectin, and with different attachment sites on plasma membranes suggests a key role for this proteoglycan in the self-assembly and insolubility of ECM components, as well as in cell adhesion and locomotion. Cleavage of the heparan sulfate (HS) chains may therefore result in degradation of the subendothelial ECM and hence may play a decisive role in extravasation of blood-borne cells. HS catabolism is observed in inflammation, wound repair, diabetes, and cancer metastasis, suggesting that enzymes which degrade HS play important roles in pathologic processes. Heparanase activity has been described in activated immune system cells and highly metastatic cancer cells (6-8), but research has been handicapped by the lack of biologic tools to explore potential causative roles of heparanase in disease conditions.

[0003] Involvement of Heparanase in Tumor Cell Invasion and Metastasis: Circulating tumor cells arrested in the capillary beds of different organs must invade the endothelial cell lining and degrade its underlying basement membrane (BM) in order to invade into the extravascular tissue(s) where they establish metastasis (9, 10). Metastatic tumor cells often attach at or near the intercellular junctions between adjacent endothelial cells. Such attachment of the metastatic cells is followed by rupture of the junctions, retraction of the endothelial cell borders and migration through the breach in the endothelium toward the exposed underlying BM (9). Once located between endothelial cells and the BM, the invading cells must degrade the subendothelial glycoproteins and proteoglycans of the BM in order to migrate out of the vascular compartment. Several cellular enzymes (e.g., collagenase IV, plasminogen activator, cathepsin B, elastase, etc.) are thought to be involved in degradation of BM (10). Among these enzymes is an endo-β-D-glucuronidase (heparanase) that cleaves HS at specific intrachain sites (6, 8, 11). Expression of a HS degrading heparanase was found to correlate with the metastatic potential of mouse lymphoma (11), fibrosarcoma and melanoma (8) cells. Moreover, elevated levels of heparanase were detected in sera from metastatic tumor bearing animals and melanoma patients (8) and in tumor biopsies of cancer patients (12).

[0004] The control of cell proliferation and tumor progression by the local microenvironment, focusing on the interaction of cells with the extracellular matrix (ECM) produced by cultured corneal and vascular endothelial cells, was investigated previously by the present inventors. This cultured ECM closely resembles the subendothelium *in vivo* in its morphological appearance and molecular composition. It contains collagens (mostly type III and IV, with smaller amounts of types I and V), proteoglycans (mostly heparan sulfate- and dermatan sulfate- proteoglycans, with smaller amounts of chondroitin sulfate proteoglycans), laminin, fibronectin,entactih and elastin (13, 14). The ability of cells to degrade HS in the cultured ECM was studied by allowing cells to interact with a metabolically sulfate labeled ECM, followed by gel filtration (Sepharose 6B) analysis of degradation products released into the culture medium (11). While intact HSPG are eluted next to the void volume of the column (Kav<0.2, Mr ~  $0.5 \times 10^6$ ), labeled degradation fragments of HS side chains are eluted more toward the V<sub>1</sub> of the column (0.5<kav<0.8, Mr =S-7x10<sup>3</sup>) (11).

[0005] The heparanase inhibitory effect of various non-anticoagulant species of heparin that might be of potential use in preventing extravasation of blood-borne cells was also investigated by the present inventors. Inhibition of heparanase was best achieved by heparin species containing 16 sugar units or more and having sulfate groups at both the N and O positions. While O-desulfation abolished the heparanase inhibiting effect of heparin, O-sulfated, N-acetylated heparin retained a high inhibitory activity, provided that the N-substituted molecules had a molecular size of about 4,000 daltons or more (7). Treatment of experimental animals with heparanase inhibitors (e.g., non-anticoagulant species of heparin) markedly reduced (>90%) the incidence of lung metastases induced by B16 melanoma, Lewis lung carcinoma and mammary adenocarcinoma cells (7, 8, 16). Heparin fractions with high and low affinity to anti-thrombin III exhibited a comparable high anti-metastatic activity, indicating that the heparanase inhibiting activity of heparin, rather than its anticoagulant activity, plays a role in the anti-metastatic properties of the polysaccharide (7).

[0006] Heparanase activity in the urine of cancer patients: In an attempt to further elucidate the involvement of heparanase in tumor progression and its relevance to human cancer, urine samples for heparanase activity were screened (16a). Heparanase activity was detected in the urine of some, but not all, cancer patients. High levels of heparanase activity were determined in the urine of patients with an aggressive metastatic disease and there was no detectable activity in the urine of healthy donors.

[0007] Heparanase activity was also found in the urine of 20% of normal and microalbuminuric insulin dependent diabetes mellitus (IDDM) patients, most likely due to diabetic nephropathy, the most important single disorder leading to renal failure in adults.

[0008] Possible involvement of heparanase in tumor angiogenesis: Fibroblast growth factors are a family of structurally related polypeptides characterized by high affinity to heparin (17). They are highly mitogenic for vascular endothelial cells and are among the most potent inducers of neovascularization (17, 18). Basic fibroblast growth factor (bFGF) has been extracted from the subendothelial ECM produced in vitro (19) and from basement membranes of the cornea (20), suggesting that ECM may serve as a reservoir for bFGF. Immunohistochemical staining revealed the localization of bFGF in basement membranes of diverse tissues and blood vessels (21). Despite the ubiquitous presence of bFGF in normal tissues, endothelial cell proliferation in these tissues is usually very low, suggesting that bFGF is somehow sequestered from its site of action. Studies on the interaction of bFGF with ECM revealed that bFGF binds to HSPG in the ECM and can be released in an active form by HS degrading enzymes (15, 20, 22). It was demonstrated that heparanase activity expressed by platelets, mast cells, neutrophils, and lymphoma cells is involved in release of active bFGF from ECM and basement membranes (23), suggesting that heparanase activity may not only function in cell migration and invasion, but may also elicit an indirect neovascular response. These results suggest that the ECM HSPG provides a natural storage depot for bFGF and possibly other heparin-binding growth promoting factors (24, 25). Displacement of bFGF from its storage within basement membranes and ECM may therefore provide a novel mechanism for induction of neovascularization in normal and pathological situations.

[0009] Recent studies indicate that heparin and HS are involved in binding of bFGF to high affinity cell surface receptors and in bFGF cell signaling (26, 27). Moreover, the size of HS required for optimal effect was similar to that of HS fragments released by heparanase (28). Similar results were obtained with vascular endothelial cells growth factor (VEGF) (29), suggesting the operation of a dual receptor mechanism involving HS in cell interaction with heparin-binding growth factors. It is therefore proposed that restriction of endothelial cell growth factors in ECM prevents their systemic action on the vascular endothelium, thus maintaining a very low rate of endothelial cells turnover and vessel growth. On the other hand, release of bFGF from storage in ECM as a complex with HS fragment, may elicit localized endothelial cell proliferation and neovascularization in processes such as wound healing, inflammation and tumor development (24, 25). [0010] Expression of heparanase by cells of the immune system: Heparanase activity correlates with the ability of activated cells of the immune system to leave the circulation and elicit both inflammatory and autoimmune responses. Interaction of platelets, granulocytes, T and B lymphocytes, macrophages and mast cells with the subendothelial ECM is associated with degradation of HS by a specific heparanase activity (6). The enzyme is released from intracellular compartments (e.g., lysosomes, specific granules, etc.) in response to various activation signals (e.g., thrombin, calcium ionophore, immune complexes, antigens, mitogens, etc.), suggesting its regulated involvement in inflammation and cellular immunity.

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## 40 Some of the observations regarding the heparanase enzyme were reviewed in reference No. 6 and are listed hereinbelow:

- [0011] First, a proteolytic activity (plasminogen activator) and heparanase participate synergistically in sequential degradation of the ECM HSPG by inflammatory leukocytes and malignant cells.
- [0012] Second, a large proportion of the platelet heparanase exists in a latent form, probably as a complex with chondroitin sulfate. The latent enzyme is activated by tumor cell-derived factor(s) and may then facilitate cell invasion through the vascular endothelium in the process of tumor metastasis.
  - [0013] Third, release of the platelet heparanase from  $\alpha$ -granules is induced by a strong stimulant (i.e., thrombin), but not in response to platelet activation on ECM.
- [0014] Fourth, the neutrophil heparanase is preferentially and readily released in response to a threshold activation and upon incubation of the cells on ECM.
  - [0015] Fifth, contact of neutrophils with ECM inhibited release of noxious enzymes (proteases, lysozyme) and oxygen radicals, but not of enzymes (heparanase, gelatinase) which may enable diapedesis. This protective role of the subendothelial ECM was observed when the cells were stimulated with soluble factors but not with phagocytosable stimulants.
  - [0016] Sixth, intracellular heparanase is secreted within minutes after exposure of T cell lines to specific antigens.
    - [0017] Seventh, mitogens (Con A, LPS) induce synthesis and secretion of heparanase by normal T and B lymphocytes maintained *in vitro*. T lymphocyte heparanase is also induced by immunization with antigen *in vivo*.
    - [0018] Eighth, heparanase activity is expressed by pre-B lymphomas and B-lymphomas, but not by plasmacytomas

and resting normal B lymphocytes.

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[0019] Ninth, heparanase activity is expressed by activated macrophages during incubation with ECM, but there was little or no release of the enzyme into the incubation medium. Similar results were obtained with human myeloid leukemia cells induced to differentiate to mature macrophages.

[0020] Tenth, T-cell mediated delayed type hypersensitivity and experimental autoimmunity are suppressed by low doses of heparanase inhibiting non-anticoagulant species of heparin (30).

[0021] Eleventh, heparanase activity expressed by platelets, neutrophils and metastatic tumor cells releases active bFGF from ECM and basement membranes. Release of bFGF from storage in ECM may elicit a localized neovascular response in processes such as wound healing, inflammation and tumor development.

[0022] Twelfth, among the breakdown products of the ECM generated by heparanase is a tri-sulfated disaccharide that can inhibit T-cell mediated inflammation *in vivo* (31). This inhibition was associated with an inhibitory effect of the disaccharide on the production of biologically active TNFα by activated T cells *in vitro* (31).

[0023] Other potential therapeutic applications: Apart from its involvement in tumor cell metastasis, inflammation and autoimmunity, mammalian heparanase may be applied to modulate: bioavailability of heparin-binding growth factors (15); cellular responses to heparin-binding growth factors (e.g., bFGF, VEGF) and cytokines (IL-8) (31 a, 29); cell interaction with plasma lipoproteins (32); cellular susceptibility to certain viral and some bacterial and protozoa infections (33, 33a, 33b); and disintegration of amyloid plaques (34). Heparanase may thus prove useful for conditions such as wound healing, angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases and viral infections. Mammalian heparanase can be used to neutralize plasma heparin; as a potential replacement of protamine. Anti-heparanase antibodies may be applied for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Common use in basic research is expected.

**[0024]** The identification of the *hpa* gene encoding for heparanase enzyme will enable the production of a recombinant enzyme in heterologous expression systems. Availability of the recombinant protein will pave the way for solving the protein structure function relationship and will provide a tool for developing new inhibitors.

[0025] Viral Infection: The presence of heparan sulfate on cell surfaces have been shown to be the principal requirement for the binding of Herpes Simplex (33) and Dengue (33a) viruses to cells and for subsequent infection of the cells. Removal of the cell surface heparan sulfate by heparanase may therefore abolish virus infection. In fact, treatment of cells with bacterial heparitinase (degrading heparan sulfate) or heparinase (degrading heparan) reduced the binding of two related animal herpes viruses to cells and rendered the cells at least partially resistant to virus infection (33). There are some indications that the cell surface heparan sulfate is also involved in HIV infection (33b).

**[0026] Neurodegenerative diseases:** Heparan sulfate proteoglycans were identified in the prion protein amyloid plaques of Genstmann-Straussler Syndrome, Creutzfeldt-Jakob disease and Scrape (34). Heparanase may disintegrate these amyloid plaques which are also thought to play a role in the pathogenesis of Alzheimer's disease.

[0027] Restenosis and Atherosclerosis: Proliferation of arterial smooth muscle cells (SMCs) in response to endothelial injury and accumulation of cholesterol rich lipoproteins are basic events in the pathogenesis of atherosclerosis and restenosis (35). Apart from its involvement in SMC proliferation (i.e., low affinity receptors for heparin-binding growth factors), HS is also involved in lipoprotein binding, retention and uptake (36). It was demonstrated that HSPG and lipoprotein lipase participate in a novel catabolic pathway that may allow substantial cellular and interstitial accumulation of cholesterol rich lipoproteins (32). The latter pathway is expected to be highly atherogenic by promoting accumulation of apoB and apoE rich lipoproteins (i.e. LDL, VLDL, chylomicrons), independent of feed back inhibition by the cellular sterol content. Removal of SMC HS by heparanase is therefore expected to inhibit both SMC proliferation and lipid accumulation and thus may halt the progression of restenosis and atherosclerosis.

[0028] There is thus a widely recognized need for, and it would be highly advantageous to have a polynucleotide encoding a polypeptide having heparanase activity, vectors including same, transduced cells expressing heparanase and a recombinant protein having heparanase activity.

## SUMMARY OF THE INVENTION

[0029] According to the present invention there is provided a polynucleotide, referred to hereinbelow as *hpa*, *hpa* cDNA or *hpa* gene, encoding a polypeptide having heparanase activity, vectors including same, transduced cells expressing heparanase and a recombinant protein having heparanase activity.

[0030] Cloning of the human hpa gene which encodes heparanase, and expression of recombinant heparanase by transfected host cells is reported.

[0031] A purified preparation of heparanase isolated from human hepatoma cells was subjected to tryptic digestion and microsequencing. The YGPDVGQPR (SEQ ID NO:8) sequence revealed was used to screen EST databases for homology to the corresponding back translated DNA sequence. Two closely related EST sequences were identified and were thereafter found to be identical. Both clones contained an insert of 1020 bp which included an open reading frame of 973 bp followed by a 27 bp of 3' untranslated region and a Poly. A tail. Translation start site was not identified.

[0032] Cloning of the missing 5' end of *hpa* was performed by PCR amplification of DNA from placenta Marathon RACE (RIM) cDNA composite using primers selected according to the EST clones sequence and the linkers of the composite. A 900 bp PCR fragment, partially overlapping with the identified 3' encoding EST clones was obtained. The joined cDNA fragment (*hpa*), 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons.

[0033] Cloning an extended. 5' sequence was enabled from the human SK-hepl cell line by PCR amplification using the Marathon RACE (RIM). The 5' extended sequence of the SK-hep 1 hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQ ID NO:9). The assembled sequence contained an open reading frame, SEQ ID NOs: 13 and 15, which encodes, as shown in SEQ ID NOs: 14 and 15, a polypeptide of 592 amino acids with, a calculated molecular weight of 66,407 daltons.

[0034] The ability of the *hpa* gene product to catalyze degradation of heparan sulfate in an *in vitro* assay was examined by expressing the entire open reading frame of *hpa* in insect cells, using the Baculovirus expression system. Extracts and conditioned media of cells infected with virus containing the *hpa* gene, demonstrated a high level of heparan sulfate degradation activity both towards soluble ECM-derived HSPG and intact ECM. This degradation activity was inhibited by heparin, which is another substrate of heparanase. Cells infected with a similar construct containing no *hpa* gene had no such activity, nor did non-infected cells. The ability of heparanase expressed from the extended 5' clone towards heparin was demonstrated in a mammalian expression system.

[0035] The expression pattern of *hpa* RNA in various tissues and cell lines was investigated using RT-PCR. It was found to be expressed only in tissues and cells previously known to have heparanase activity.

[0036] A panel of monochromosomal human/CHO and human/mouse somatic cell hybrids was used to localize the human heparanase gene to human chromosome 4. The newly isolated heparanase sequence can be used to identify a chromosome region harboring a human heparanase gene in a chromosome spread.

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[0037] According to further features in preferred embodiments of the invention described below, there is provided a polynucleotide fragment which includes a polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

[0038] According to still further features in the described preferred embodiments the polynucleotide fragment includes nucleotides 63-1691 of SEQ ID NO:9 or nucleotides 139-1869 of SEQ ID. NO:13, which encode the entire human heparanase enzyme.

[0039] According to still further features in the described preferred embodiments there is provided a polynucleotide fragment which includes a polynucleotide sequence capable of hybridizing with hpa cDNA, especially with nucleotides 1-721 of SEQ ID NO:9.

[0040] According to still further features in the described preferred embodiments the polynucleotide sequence which encodes the polypeptide having heparanase activity shares at least 60 % homology, preferably at least 70 % homology, more preferably at least 80 % homology, most preferably at least 90 % homology with SEQ ID NOs:9 or 13.

[0041] According to still further features in the described preferred embodiments the polynucleotide fragment according to the present invention includes a portion (fragment) of SEQ ID NOs:9, or 13. For example, such fragments could include nucleotides 63-721 of SEQ ID NO:9 and/or a segment of SEQ ID NO:9 which encodes a polypeptide having the heparanase catalytic activity.

[0042] According to still further features in the described preferred embodiments the polypeptide encoded by the polynucleotide fragment includes an amino acid sequence as set forth in SEQ ID NOs:10 or 14 or a functional part thereof. [0043] According to still further features in the described preferred embodiments the polynucleotide sequence encodes a polypeptide having heparanase activity, which shares at least 60 % homology, preferably at least 70 % homology, more preferably at least 80 % homology, most preferably at least 90 % homology with SEQ ID NOs:10 or 14.

[0044] According to still further features in the described preferred embodiments the polynucleotide fragment encodes a polypeptide having heparanase activity, which may therefore be allelic, species and/or induced variant of the amino acid sequence set forth in SEQ ID NOs:10 or 14. It is understood that any such variant may also be considered a homolog. [0045] According to still further features in the described preferred embodiments there is provided a single stranded polynucleotide fragment which includes a polynucleotide sequence complementary to at least a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity as described above.

[0046] According to still further features in the described preferred embodiments there is provided a vector including a polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

[0047] The vector may be of any suitable type including but not limited to a phage, virus, plasmid, phagemid, cosmid, bacmid or even an artificial chromosome. The polynucleotide sequence encoding a polypeptide having heparanase catalytic activity may include any of the above described polynucleotide fragments.

[0048] According to still further features in the described preferred embodiments there is provided a host cell which includes an exogenous polynucleotide fragment including a polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

[0049] The exogenous polynucleotide fragment may be any of the above described fragments. The host cell may be

of any type such as prokaryotic cell, eukaryotic cell, a cell line, or a cell as a portion of a multicellular organism (e.g., cells of a transgenic organism).

[0050] According to still further features in the described preferred embodiments there is provided a recombinant protein including a polypeptide having heparanase catalytic activity.

[0051] According to still further features in the described preferred embodiments there is provided a pharmaceutical composition comprising as an active ingredient a recombinant protein having heparanase catalytic activity.

[0052] According to still further features in the described preferred embodiments there is provided a medical equipment comprising a medical device containing, as an active ingredient a recombinant protein having heparanase catalytic activity.

[0053] According to still further features in the described preferred embodiments there is provided a heparanase overexpression system comprising a cell overexpressing heparanase catalytic activity.

[0054] According to still further features in the described preferred embodiments there is provided a method of identifying a chromosome region harboring a human heparanase gene in a chromosome spread comprising the steps of (a) hybridizing the chromosome spread with a tagged polynucleotyde probe encoding heparanase; (b) washing the chromosome spread, thereby removing excess of non-hybridized probe; and (c) searching for signals associated with said hybridized tagged polynucleotyde probe, wherein detected signals being indicative of a chromosome region harboring a human heparanase gene.

**[0055]** The present invention can be used to develop new drugs to inhibit tumor cell metastasis, inflammation and autoimmunity. The identification of the *hpa* gene encoding for heparanase enzyme enables the production of a recombinant enzyme in heterologous expression systems.

## BRIEF DESCRIPTION OF THE DRAWINGS

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[0056] The invention herein described, by way of example only, with reference to the accompanying drawings, wherein:

FIG. 1 presents nucleotide sequence and deduced amino acid sequence of *hpa* cDNA. A single nucleotide difference at position 799 (A to T) between the EST (Expressed Sequence Tag) and the PCR amplified cDNA (reverse transcribed RNA) and the resulting amino acid substitution (Tyr to Phe) are indicated above and below the substituted unit, respectively. Cysteine residues and the poly adenylation consensus sequence are underlined. The asterisk denotes the stop codon TGA.

FIG. 2 demonstrates degradation of soluble sulfate labeled HSPG substrate by lysates of High Five (RIM) cells infected with pFhpa2 virus. Lysates of High Five (RIM) cells that were infected with pFhpa2 virus (•) or control pF2 virus (□) were incubated (18 h, 37 °C) with sulfate labeled ECM-derivrd soluble HSPG (peak I). The incubation medium was then subjected to gel filtration on Sepharose 6B. Low molecular weight HS degradation fragments (peak II) were produced only during incubation with the pFhpa2 infected cells, but there was no degradation of the HSPG substrate (♦) by lysates of pF2 infected cells.

FIGs. 3a-b demonstrate degradation of soluble sulfate labeled HSPG substrate by the culture medium of pFhpa2 and pFhpa4 infected cells. Culture media of High Five (RIM) cells infected with pFhpa2 (3a) or pFhpa4 (3b) viruses (•), or with control viruses (□) were incubated (18 h, 37 °C) with sulfate labeled ECM-derived soluble HSPG (peak I, ♦). The incubation media were then subjected to gel filtration on Sepharose 6B. Low molecular weight HS degradation fragments (peak II) were produced only during incubation with the hpa gene containing viruses. There was no degradation of the HSPG substrate by the culture medium of cells infected with control viruses.

FIG. 4 presents size fractionation of heparanase activity expressed by pFhpa2 infected cells. Culture medium of pFhpa2 infected High Five (RIM) cells was applied onto a 50 kDa cut-off membrane. Heparanase activity (conversion of the peak I substrate, (�) into peak II HS degradation fragments) was found in the high (> 50 kDa) (\*), but not low (< 50 kDa) (o) molecular weight compartment.

FIGs. 5a-b demonstrate the effect of heparin on heparanase activity expressed by pFhpa2 and pFhpa4 infected High Five (RIM) cells. Culture media of pFhpa2 (5a) and pFhpa4 (5b) infected High Five (RIM) cells were incubeted (18 h, 37 °C) with sulfate labeled ECM-derived soluble HSPG (peak.I,  $\spadesuit$ ) in the absence ( $\bullet$ ) or presence ( $\triangle$ ) of 10  $\mu$ g/ml heparin. Production of low molecular weight HS degradation fragments was completely abolished in the presence of heparin, a potent inhibitor of heparanase activity (6, 7).

FIGs. 6a-b demonstrate degradation of sulfate labeled intact ECM by virus infected High Five (RIM) and Sf21 Cells. High Five (RIM) (6a) and Sf21 (6b) cells were plated on sulfate labeled ECM and infected (48 h, 28 °C) with pFhpa4 (•) or control pF1 (□) viruses. Control non-infected Sf21 cells (R) were plated on the labeled ECM as well. The pH of the cultured medium was adjusted to 6.0 - 6.2 followed by 24 h incubation at 37 °C. Sulfate labeled material released into the incubation medium was analyzed by gel filtration on Sepharose 6B. HS degradation fragments were produced only by cells infected with the *hpa* containing virus.

FIG. 7a-b demonstrate degradation of sulfate labeled intact ECM by virus infected cells. High Five (RIM) (7a) and

Sf21 (7b) cells were plated on sulfate labeled ECM and infected (48 h, 28 °C)with pFhpa4 (•) or control pF1 (□) viruses. Control non-infected Sf21 cells (R) were plate on labeled ECM as well. The pH of the cultured medium was adjusted to 6.0 - 6.2, followed by 48 h incubation at 28 °C. Sulfate labeled degradation fragments released into the incubation medium was analyzed by gel filtration on Sepharose 6B. HS degradation fragments were produced only by cells infected with the hpa containing virus.

FIGs. 8a-b demonstrate degradation of sulfate labeled intact ECM by the culture medium of pFhpa4 infected cells. Culture media of High Five (RIM) (8a) and Sf21 (8b) cells that were infected with pFhpa4 (•) or control pF1 (□) viruses were incubated (48 h, 37 °C, pH 6.0) with intact sulfate labeled ECM. The ECM was also incubated with the culture medium of control non-infected Sf21 cells (R). Sulfate labeled material released into the reaction mixture was subjected to gel filtration analysis. Heparanase activity was detected only in the culture medium of pFhpa4 infected cells..

FIGs. 9a-b demonstrate the effect of heparin on heparanase activity in the culture medium of pF*hpa*4 infected cells. Sulfate labeled ECM was incubated (24 h, 37 °C, pH 6.0) with culture medium of pF*hpa*4 infected High Five (RIM) ·(9a) and Sf2 (9b) cells in the absence (•) or presence (V) of 10 μg/ml heparin. Sulfate labeled material released into the incubation medium was subjected to gel filtration on Sepharose 6B. Heparanase activity (production of peak II HS degradation fragments) was completely inhibited in the presence of heparin.

FIGS. 10a-b demonstrate purification of recombinant heparanase on heparin-Sepharose. Culture medium of Sf21 cells infected with pFhpa4 virus was subjected to heparin-Sepharose chromatography. Elution of fractions was performed with 0.3 - 2 M NaCl gradient (�). Heparanase activity in the eluted fractions is demonstrated in Figure 10a (\*). Fractions 15-28 were subjected to 15% SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining. A correlation is demonstrated between a major protein band (MW ~ 63,000) in fractions 19-24 and heparanase activity.

FIGs. 11a-b demonstrate purification of recombinant heparanase on a Superdex 75 gel filtration column. Active fractions eluted from heparin-Sepharose (Figure 10a) were pooled, concentrated and applied onto Superdex 75 FPLC column. Fractions were collected and aliquots of each fraction were tested for heparanase activity (C, Figure 11a) and analyzed by SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining (Figure 11b). A correlation is seen between the appearance of a major protein band (MW ~ 63,000) in fractions 4 - 7 and heparanase activity.

FIGs. 12a-e demonstrate expression of the *hpa* gene by RT-PCR with total RNA from human embryonal tissues (12a), human extra-embryonal tissues (12b) and cell lines from different origins (12c-e). RT-PCR products using *hpa* specific primers (I), primers for GAPDH housekeeping gene (II), and control reactions without reverse transcriptase demonstrating absence of genomic DNA or other contamination in RNA samples (III). M- DNA molecular weight marker VI (Boehringer Mannheim). For 12a: lane 1 - neutrophil cells (adult), lane 2 - muscle, lane 3 - thymus, lane 4 - heart, lane 5 - adrenal. For 12b: lane 1 - kidney, lane 2 - placenta (8 weeks), lane 3 - placenta (11 weeks), lanes 4-7 - mole (complete hydatidiform mole), lane 8 - cytotrophoblast cells (freshly isolated), lane 9 - cytotrophoblast cells (1.5 h *in vitro*), lane 10 - cytotrophoblast cells (6 h *in vitro*), lane 11 - cytotrophoblast cells (18 h *in vitro*), lane 12 - cytotrophoblast cells (48 h *in vitro*). For 12c: lane 1 - JAR bladder cell line, lane 2 - NCITT testicular tumor cell line, lane 3 - SW-480 human hepatoma cell line, lane 4 - HTR (cytotrophoblasts transformed by SV40), lane 5 - HPTLP-1 hepatocellular carcinoma cell line, lane 6 - EJ-28 bladder carcinoma cell line. For 12d: lane 1 - SK-hep-1 human hepatoma cell line, lane 2 - DAMI human megakaryocytic cell line, lane 3 - DAMI cell line + PMA, lane 4 - CHRF cell line + PMA, lane 5 - CHRF cell line. For 12e: lane 1 - ABAE bovine aortic endothelial cells, lane 2 - 1063 human ovarian cell line, lane 3 - human breast carcinoma MDA435 cell line, lane 4 - human breast carcinoma MDA431 cell line.

FIG. 13 presents a comparison between nucleotide sequences of the human *hpa* and a mouse EST cDNA fragment (SEQ ID NO:12) which is 80 % homologous to the 3' end (starting at nucleotide 1066 of SEQ ID NO:9) of the human *hpa*. The aligned termination codons are underlined.

FIG. 14 demonstrates the chromosomal localization of the *hpa* gene. PCR products of DNA derived from somatic cell hybrids and of genomic DNA of hamster, mouse and human of were separated on 0.7 % agarose gel following amplification with *hpa* specific primers. Lane 1 - Lambda DNA digested with *Bst*EII, lane 2 - no DNA control, lanes 3 - 29, PCR amplification products. Lanes 3-5 - human, mouse and hamster genomic DNA, respectively. Lanes 6-29, human monochromosomal somatic cell hybrids representing chromosomes 1-22 and X and Y, respectively. Lane 30 - Lambda DNA digested with *Bst*EII. An amplification product of approximately 2.8 Kb is observed only in lanes 5 and 9, representing human genomic DNA and DNA derived from cell hybrid carrying human chromosome 4, respectively. These results demonstrate that the *hpa* gene is localized in human chromosome 4.

## **DESCRIPTION OF THE PREFERRED EMBODIMENTS**

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[0057] The present invention is of a polynucleotide, referred to hereinbelow interchangeably as hpa, hpa cDNA or

hpa gene, encoding a polypeptide having heparanase activity, vectors including same, transduced cells expressing heparanase and a recombinant protein having heparanase activity.

[0058] Before explaining at least one embodiment of the invention in detail, it is to be understood that the invention is not limited in its application to the details of construction and the arrangement of the components set forth in the following description or illustrated in the drawings.

[0059] Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

[0060] The present invention can be used to develop treatments for various diseases, to develop diagnostic assays for these diseases and to provide new tools for basic research especially in the fields of medicine and biology.

[0061] Specifically, the present invention can be used to develop new drugs to inhibit tumor cell metastasis, inflammation and autoimmunity. The identification of the *hpa* gene encoding for the heparanase enzyme enables the production of a recombinant enzyme in heterologous expression systems.

[0062] Furthermore, the present invention can be used to modulate bioavailability of heparin-binding growth factors, cellular responses to heparin-binding growth factors (e.g., bFGF, VEGF) and cytokines (IL-8), cell interaction with plasma lipoproteins, cellular susceptibility to viral, protozoa and some bacterial infections, and disintegration of neurodegenerative plaques. Recombinant heparanase is thus a potential treatment for wound healing, angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases (such as, for example, Genstmann-Straussler Syndrome, Creutzfeldt-Jakob disease, Scrape and Alzheimer's disease) and certain viral and some bacterial and protozoa infections. Recombinant heparanase can be used to neutralize plasma heparin, as a potential replacement of protamine.

[0063] As used herein, the term "modulate" includes substantially inhibiting, slowing or reversing the progression of a disease, substantially ameliorating clinical symptoms of a disease or condition, or substantially preventing the appearance of clinical symptoms of a disease or condition. A "modulator" therefore includes an agent which may modulate a disease or condition. Modulation of viral, protozoa and bacterial infections includes any effect which substantially interrupts, prevents or reduces any viral, bacterial or protozoa activity and/or stage of the virus, bacterium or protozoon life cycle, or which reduces or prevents infection by the virus, bacterium or protozoon in a subject, such as a human or lower animal

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[0064] As used herein, the term "wound" includes any injury to any portion of the body of a subject including, but not limited to, acute conditions such as thermal burns, chemical burns, radiation burns, burns caused by excess exposure to ultraviolet radiation such as sunburn, damage to bodily tissues such as the perineum as a result of labor and childbirth, including injuries sustained during medical procedures such as episiotomies, trauma-induced injuries including cuts, those injuries sustained in automobile and other mechanical accidents, and those caused by bullets, knives and other weapons, and post-surgical injuries, as well as chronic conditions such as pressure sores, bedsores, conditions related to diabetes and poor circulation, and all types of acne, etc.

[0065] Anti-heparanase antibodies, raised against the recombinant enzyme, would be useful for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Such antibodies may also serve as neutralizing agents for heparanase activity.

**[0066]** Cloning of the human *hpa* gene encoding heparanase and expressing recombinant heparanase by transfected cells is herein reported. This is the first mammalian heparanase gene to be cloned.

[0067] A purified preparation of heparanase isolated from human hepatoina cells was subjected to tryptic digestion and microsequencing.

[0068] The YGPDVGQPR (SEQ ID NO:8) sequence revealed was used to screen EST databases for homology to the corresponding back translated DNA sequences. Two closely related EST sequences were identified and were thereafter found to be identical.

[0069] Both clones contained an insert of 1020 bp which includes an open reading frame of 973 bp followed by a 3' untranslated region of 27 bp and a Poly A tail, whereas a translation start site was not identified.

[0070] Cloning of the missing 5' end was performed by PCR amplification of DNA from placenta marathon RACE (RIM) cDNA composite using primers selected according to the EST clones sequence and the linkers of the composite. [0071] A 900 bp PCR fragment, partially overlapping with the identified 3' encoding EST clones was obtained. The joined cDNA fragment (hpa), 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes, as shown in Figure 1 and SEQ ID NO:11, a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons.

[0072] A single nucleotide difference at position 799 (A to T) between the EST clones and the PCR amplified cDNA was observed. This difference results in a single amino acid substitution (Tyr to Phe) (Figure 1). Furthermore, the published EST sequences contained an unidentified nucleotide, which following DNA sequencing of both the EST clones was resolved into two nucleotides (G and C at positions 1630 and 1631 in SEQ ID NO:9, respectively).

**[0073]** The ability of the *hpa* gene product to catalyze degradation of heparan sulfate in an *in vitro* assay was examined by expressing the entire open reading frame in insect cells, using the Baculovirus expression system.

[0074] Extracts and conditioned media of cells infected with virus containing the hpa gene, demonstrated a high level

of heparan sulfate degradation activity both towards soluble ECM-derived HSPG and intact ECM, which was inhibited by heparin, while cells infected with a similar construct containing no *hpa* gene had no such activity, nor did non-infected cells.

[0075] The expression pattern of hpa RNA in various tissues and cell lines was investigated using RT-PCR. It was found to be expressed only in tissues and cells previously known to have heparanase activity.

[0076] Cloning an extended 5' sequence was enabled from the human SK-hep1 cell line by PCR amplification using the Marathon RACE (RIM) The 5' extended sequence of the SK-hep1 hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQ ID NO:9). The assembled sequence contained an open reading frame, SEQ ID NOs: 13 and 15, which encodes, as shown in SEQ ID NOs:14 and 15, a polypeptide of 592 amino acids, with a calculated molecular weight of 66,407 daltons. This open reading frame was shown to direct the expression of catalitically active heparanase in a mammalian cell expression system. The expressed heparanase was detectable by anti heparanase antibodies in Western blot analysis.

[0077] A panel of monochromosomal human/CHO and human/mouse somatic cell hybrids was used to localize the human heparanase gene to human chromosome 4. The newly isolated heparanase sequence can therefore be used to identify a chromosome region harboring a human heparanase gene in a chromosome spread.

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[0078] Thus, according to the present invention there is provided a polynucleotide fragment (either DNA or RNA, either single stranded or double stranded) which includes a polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

[0079] The term "heparanase catalytic activity" or its equivalent term "heparanase activity" both refer to a mammalian endoglycosidase hydrolyzing activity which is specific for heparan or heparan sulfate proteoglycan substrates, as opposed to the activity of bacterial enzymes (heparinase I, II and III) which degrade heparin or heparan sulfate by means of β-elimination (37).

[0080] In a preferred embodiment of the invention the polynucleotide fragment includes nucleotides 63-1691 of SEQ ID NO:9, or nucleotides 139-1869 of SEQ ID NO: 13, which encode the entire human heparanase enzyme.

[0081] However, the scope of the present invention is not limited to human heparanase since this is the first disclosure of an open reading frame (ORF) encoding any mammalian heparanase. Using the *hpa* cDNA, parts thereof or synthetic oligonucleotides designed according to its sequence will enable one ordinarily skilled in the art to identify genomic and/or cDNA clones including homologous sequences from other mammalian species.

[0082] The present invention is therefore further directed at a polynucleotide fragment which includes a polynucleotide sequence capable of hybridizing (base pairing under either stringent or permissive hybridization conditions, as for example described in Sambrook, J.; Fritsch, E.F., Maniatis, T. (1989) Molecular Cloning. A Laboratory Manual. Cold Spring Harbor Laboratory Press, New York.) with hpa cDNA, especially with nucleotides 1-721 of SEQ ID NO:9.

[0083] In fact, any polynucleotide sequence which encodes a polypeptide having heparanase activity and which shares at least 60 % homology, preferably at least 70 % homology, more preferably at least 80 % homology, most preferably at least 90 % homology with SEQ ID NOs:9 or 13 is within the scope of the present invention.

[0084] The polynucleotide fragment according to the present invention may include any part of SEQ ID NOs: 9 or 13. For example, it may include nucleotides 63-721 of SEQ ID NO:9, which is a novel sequence. However, it may include any segment of SEQ ID NOs:9 or 13 which encodes a polypeptide having the heparanase catalytic activity.

[0085] When the phrase "encodes a polypeptide having heparanase catalytic activity" is used herein and in the claims section below it refers to the ability of directing the synthesis of a polypeptide which, if so required for its activity, following post translational modifications, such as but not limited to, proteolysis (e.g., removal of a signal peptide and of a pro- or preprotein sequence), methionine modification, glycosylation, alkylation (e.g., methylation), acetylation, etc., is catalytically active in degradation of, for example, ECM and cell surface associated HS.

[0086] In a preferred embodiment of the invention the polypeptide encoded by the polynucleotide fragment includes an amino acid sequence as set forth in SEQ ID NOs:10 or 14 or a functional part thereof, i.e., a portion harboring heparanase catalytic activity.

[0087] However, any polynucleotide fragment which encodes a polypeptide having heparanase activity is within the scope of the present invention. Therefore, the polypeptide may be allelic, species and/or induced variant of the amino acid sequence set forth in SEQ ID NOs: 10 or 14 or functional part thereof.

[0088] In fact, any polynucleotide sequence which encodes a polypeptide having heparanase activity, which shares at least 60 % homology, preferably at least 70 % homology, more preferably at least 80 % homology, most preferably at least 90 % homology with SEQ ID NOs:10 or 14 is within the scope of the present invention.

[0089] The invention is also directed at providing a single stranded polynucleotide fragment which includes a polynucleotide sequence complementary to at least a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity as described above. The term "complementary" as used herein refers to the ability of base pairing. [0090] The single stranded polynucleotide fragment may be DNA or RNA or even include nucleotide analogs (e.g., thioated nucleotides), it may be a synthetic oligonucleotide or manufactured by transduced host cells, it may be of any desired length which still provides specific base pairing (e.g., 8 or 10, preferably more, nucleotides long) and it may

include mismatches that do not hamper base pairing.

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[0091] The invention is further directed at providing a vector which includes a polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

[0092] The vector may be of any type. It may be a phage which infects bacteria or a virus which infects eukaryotic cells. It may also be a plasmid, phagemid, cosmid, bacmid or an artificial chromosome. The polynucleotide sequence encoding a polypeptide having heparanase catalytic activity may include any of the above described polynucleotide fragments.

[0093] The invention is further directed at providing a host cell which includes an exogenous polynucleotide fragment encoding a polypeptide having heparanase catalytic activity.

[0094] The exogenous polynucleotide fragment may be any of the above described fragments. The host cell may be of any type. It may be a prokaryotic cell, an eukaryotic cell, a cell line, or a cell as a portion of an organism. The exogenous polynucleotide fragment may be permanently or transiently present in the cell. In other words, transduced cells obtained following stable or transient transfection, transformation or transduction are all within the scope of the present invention. The term "exogenous" as used herein refers to the fact that the polynucleotide fragment is externally introduced into the cell. Therein it may be present in a single of any number of copies, it may be integrated into one or more chromosomes at any location or be present as an extrachromosomal material.

[0095] The invention is further directed at providing a heparanase overexpression system which includes a cell overexpressing heparanase catalytic activity. The cell may be a host cell transiently or stably transfected or transformed with any suitable vector which includes a polynucleotide sequence encoding a polypeptide having heparanase activity and a suitable promoter and enhancer sequences to direct overexpression of heparanase. However, the overexpressing cell may also be a product of an insertion (e.g., via homologous recombination) of a promoter and/or enhancer sequence upstream to the endogenous heparanase gene of the expressing cell, which will direct overexpression from the endogenous gene. The term "overexpression" as used herein in the specification and claims below refers to a level of expression which is higher than a basal level of expression typically characterizing a given cell under otherwise identical conditions.

[0096] The invention is further directed at providing a recombinant protein including a polypeptide having heparanase

[0097] The recombinant protein may be purified by any conventional protein purification procedure close to homogeneity and/or be mixed with additives. The recombinant protein may be manufactured using any of the cells described above. The recombinant protein may be in any form. It may be in a crystallized form, a dehydrated powder form or in solution. The recombinant protein may be useful in obtaining pure heparanase, which in turn may be useful in eliciting anti-heparanase antibodies, either poly or monoclonal antibodies, and as a screening active ingredient in an anti-heparanase inhibitors or drugs screening assay or system.

[0098] The invention is further directed at providing a pharmaceutical composition which include as an active ingredient a recombinant protein having heparanase catalytic activity.

[0099] Formulations for topical administration may include, but are not limited to, lotions, ointments, gels, creams, suppositories, drops, liquids, sprays and powders. Conventional pharmaceutical carriers, aqueous, powder or oily bases, thickeners and the like may be necessary or desirable. Coated condoms, stents, active pads, and other medical devices may also be useful. In fact the scope of the present invention includes any medical equipment such as a medical device containing, as an active ingredient, a recombinant protein having heparanase catalytic activity.

[0100] Compositions for oral administration include powders or granules, suspensions or solutions in water or non-aqueous media, sachets, capsules or tablets. Thickeners, diluents, flavorings, dispersing aids, emulsifiers or binders may be desirable.

[0101] Formulations for parenteral administration may include, but are not limited to; sterile aqueous solutions which may also contain buffers, diluents and other suitable additives.

[0102] Dosing is dependent on severity and responsiveness of the condition to be treated, but will normally be one or more doses per day, with course of treatment lasting from several days to several months or until a cure is effected or a diminution of disease state is achieved. Persons ordinarily skilled in the art can easily determine optimum dosages, dosing methodologies and repetition rates.

**[0103]** Further according to the present invention there is provided a method of identifying a chromosome region harboring a human heparanase gene in a chromosome spread, the method is executed implementing the following method steps, in which in a first step the chromosome spread (either interphase or metaphase spread) is hybridized with a tagged polynucleotyde probe encoding heparanase. The tag is preferably a fluorescent tag. In a second step according to the method the chromosome spread is washed, thereby excess of non-hybridized probe is removed. Finally, signals associated with the hybridized tagged polynucleotyde probe are searched for, wherein detected signals being indicative of a chromosome region harboring the human heparanase gene. One ordinarily skilled in the art would know how to use the sequences disclosed herein in suitable labeling reactions and how to use the tagged probes to detect, using *in situ* hybridization, a chromosome region harboring a human heparanase gene.

[0104] Reference is now made to the following examples, which together with the above descriptions, illustrate the

invention in a non-limiting fashion.

#### **EXAMPLES**

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[0105] The following protocols and experimental details are referenced in the Examples that follow:

[0106] Purification and characterization of heparanase from a human hepatoma cell line and human placenta: A human hepatoma cell line (Sk-hep-1) was chosen as a source for purification of a human tumor-derived heparanase. Purification was essentially as described in U.S. Pat. No. 5,362,641 to Fuks.

[0107] Briefly, 500 liter, 5x10<sup>11</sup> cells were grown in suspension and the heparanase enzyme was purified about 240,000 fold by applying the following steps: (i) cation exchange (CM-Sephadex) chromatography performed at pH 6.0, 0.3-1.4 M NaCl gradient; (ii) cation exchange (CM-Sephadex) chromatography performed at pH 7.4 in the presence of 0.1.% CHAPS, 0.3-1.1 M NaCl gradient; (iii) heparin-Sepharose chromatography performed at pH 7.4 in the presence of 0.1% CHAPS and 1 M NaCl gradient; (iv) ConA-Sepharose chromatography performed at pH 6.0 in buffer containing 0.1 % CHAPS and 1 M NaCl, elution with 0.25 M α-methyl mannoside; and (v) HPLC cation exchange (Mono-S) chromatography performed at pH 7.4 in the presence of 0.1 % CHAPS, 0.25-1 M NaCl gradient.

[0108] Active fractions were pooled, precipitated with TCA and the precipitate subjected to SDS polyacrylamide gel electrophoresis and/or tryptic digestion and reverse phase HPLC. Tryptic peptides of the purified protein were separated by reverse phase HPLC (C8 column) and homogeneous peaks were subjected to amino acid sequence analysis.

[0109] The purified enzyme was applied to reverse phase HPLC and subjected to N-terminal amino acid sequencing using the amino acid sequencer (Applied Biosystems).

[0110] Cells: Cultures of bovine corneal endothelial cells (BCECs) were established from steer eyes as previously described (19, 38). Stock cultures were maintained in DMEM (1 g glucose/liter) supplemented with 10 % newborn calf serum and 5 % FCS. bFGF (1 ng/ml) was added every other day during the phase of active cell growth (13,14).

[0111] Preparation of dishes coated with ECM: BCECs (second to fifth passage) were plated into 4-well plates at an initial density of 2 x 10<sup>5</sup> cells/ml, and cultured in sulfate-free Fisher medium plus 5 % dextran T-40 for 12 days Na<sub>2</sub><sup>35</sup>SO<sub>4</sub> (25 μCi/ml) was added on day 1 and 5 after seeding and the cultures were incubated with the label without medium change. The subendothelial ECM was exposed by dissolving (5 min., room temperature) the cell layer with PBS containing 0.5 % Triton. X-100 and 20 mM NH<sub>4</sub>OH, followed by four washes with PBS. The ECM remained intact; free of cellular debris and firmly attached to the entire area of the tissue culture dish (19, 22).

[0112] To prepare soluble sulfate labeled proteoglycans (peak I material), the ECM was digested with trypsin (25 µg/ml, 6 h, 37 °C), the digest was concentrated by reverse dialysis and the concentrated material was applied onto a Sepharose 6B gel filtration column. The resulting high molecular weight material (Kav< 0.2, peak I) was collected. More than 80 % of the labeled material was shown to be composed ofheparan sulfate proteoglycans (11, 39).

[0113] Heparanase activity: Cells (1 x  $10^6/35$ -mm dish), cell lysates or conditioned media were incubated on top of  $^{35}$ S-labeled ECM (18 h, 37 °C) in the presence of 20 mM phosphate buffer (pH 6.2). Cell lysates and conditioned media were also incubated with sulfate labeled peak I material ( $10\text{-}20~\mu\text{I}$ ). The incubation medium was collected, centrifuged (18,000~x g, 4 °C, 3 min.), and sulfate labeled material analyzed by gel filtration on a Sepharose CL-6B column (0.9~x 30 cm). Fractions (0.2 ml) were eluted with PBS at a flow rate of 5 ml/h and counted for radioactivity using Bio-fluor scintillation fluid. The excluded volume ( $V_0$ ) was marked by blue dextran and the total included volume ( $V_1$ ) by phenol red. The latter was shown to comigrate with free sulfate (7, 1, 23). Degradation fragments of HS side chains were eluted from Sepharose 6B at 0.5 < Kav < 0.8 (peak II) (7, 11, 23). A nearly intact HSPG released from ECM by trypsin - and, to a lower extent, during incubation with PBS alone - was eluted next to  $V_0$  (Kav < 0.2, peak I). Recoveries of labeled material applied on the columns ranged from 85 to 95 % in different experiments (11). Each experiment was performed at least three times and the variation of elution positions (Kav values) did not exceed +/- 15 %.

[0114] Cloning of hpa cDNA: cDNA clones 257548 and 260138 were obtained from the I.M.A.G.E Consortium (2130 Memorial Parkway SW, Hunstville, AL 35801). The cDNAs were originally cloned in EcoRI and Notl cloning sites in the plasmid vector pT3T7D-Pac. Although these clones are reported to be somewhat different, DNA sequencing demonstrated that these clones are identical to one another. Marathon RACE (RIM) (rapid amplification of cDNA ends) human placenta (poly-A) cDNA composite was a gift of Prof Yossi Shiloh of Tel Aviv University. This composite is vector free, as it includes reverse transcribed cDNA fragments to which double, partially single stranded adapters are attached on both sides. The construction of the specific composite employed is described in reference 39a.

[0115] Amplification of hp3 PCR fragment was performed according to the protocol provided by Clontech laboratories. The template used for amplification was a sample taken from the above composite. The primers used for amplification were:

First step: 5'-primer: AP1: 5'-CCATCCTAATACGACTCACTATAGGG C-3', SEQ ID NO:1; 3'-primer: HPL229: 5'-GTAGTGATGCCATGTAACTGA ATC-3', SEQ ID NO:2.

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[0116] Second step: nested 5'-primer AP2: 5'-ACTCACTATAGGGCTCGAGCG GC-3', SEQ ID NO:3; nested 3'-primer: HPL171: 5'-GCATCTTAGCCGTCT TTCTTCG-3', SEQ ID NO:4. The HPL229 and HPL 171 were selected according to the sequence of the EST clones. They include nucleotides 933-956 and 876-897 of SEQ ID NO:9, respectively. [0117] PCR program was 94 °C - 4 min., followed by 30 cycles of 94 °C - 40 sec., 62 °C - 1 min., 72°C - 2.5 min. Amplification was performed with Expand High Fidelity (Boehringer Mannheim). The resulting ca. 900 bp hp3 PCR product was, digested with *Bfn* and *Pvul*I. Clone 257548 (phpa1) was digested with *Eco*RI, followed by end filling and was then further digested with *Bfn*. Thereafter the *Pvul*I - *Bfn* fragment of the hp3 PCR product was cloned into the blunt end - *Bfn* end of clone phpa1 which resulted in having the entire cDNA cloned in pT3T7-pac vector, designated phpa2. [0118] *DNA Sequencing:* Sequence determinations were performed with vector specific and gene specific primers, using an automated DNA sequencer (Applied Biosystems, model 373A). Each nucleotide was read from at least two independent primers.

[0119] Computer analysis of sequences: Database searches for sequence similarities were performed using the Blast network service. Sequence analysis and alignment of DNA and protein sequences were done using the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the University of Wisconsin, [0120] RT-PCR: RNA was prepared using TRI-Reagent (Molecular research center Inc.) according to the manufacturer instructions. 1.25 µg were taken for reverse transcription reaction using MuMLV Reverse transcriptase (Gibco BRL) and Oligo (dT)<sub>15</sub> primer, SEQ ID NO:5, (Promega). Amplification of the resultant first strand cDNA was performed with Taq polymerase (Promega). The following primers were used:

HPU-355: 5'-TTCGATCCCAAGAAGGAATCAAC-3', SEQ ID NO:6, nucleotides 372-394 in SEQ ID NO:9 or 11. HPL-229: 5'-GTAGTGATGCCATGTAACTGAATC-3', SEQ ID NO:7, nucleotides 933-956 in SEQ ID NO:9 or 11. PCR program: 94 °C - 4 min., followed by 30 cycles of 94 °C - 40 sec., 62 °C - I min., 72 °C - I min.

[0121] Expression of recombinant heparanase in insect cells; Cells, High Five (RIM) and Sf21 insect cell lines were maintained as monolayer cultures in SF900II-SFM medium (GibcoBRL).

[0122] Recombinant Baculovirus: Recombinant virus containing the hpa gene was constructed using the Bac to Bac system (GibcoBRL). The transfer vector pFastBac was digested with Sall and Nofl and ligated with a 1-7 kb fragment of phpa2 digested with Xhol and Nofl. The resulting plasmid was designated pFasthpa2. An identical plasmid designated pFasthpa4 was prepared as a duplicate and both independently served for further experimentations. Recombinant bacmid was generated according to the instructions of the manufacturer with pFasthpa2, pFasthpa4 and with pFastBac. The latter served as a negative control. Recombinant bacmid DNAs were transfected into Sf21 insect cells. Five days after transfection recombinant viruses were harvested and used to infect High Five (RIM) cells. 3 x 10<sup>6</sup> cells in T-25 flasks. Cells were harvested 2 - 3 days after infection. 4 x 10<sup>6</sup> cells were centrifuged and resuspended in a reaction buffer containing 20 mM phosphate citrate buffer, 50 mM NaCl. Cells underwent three cycles of freeze and thaw and lysates were stored at -80 °C. Conditioned medium was stored at 4 °C.

[0123] Partial purification of recombinant heparanase: Partial purification of recombinant heparanase was performed by heparin-Sepharose column chromatography followed by Superdex 75 column gel filtration. Culture medium (150 ml) of Sf21 cells infected with pFhpa4 virus was subjected to heparin-Sepharose chromatography. Elution of 1 ml fractions was performed with 0.35 - 2 M NaCl gradient in presence of 0.1 % CHAPS and 1 mM DTT in 10 mM sodium acetate buffer, pH 5.0. A 25 µl sample of each fraction was tested for heparanase activity. Heparanase activity was eluted at the range of 0.65 - 1.1 M NaCl (fractions 18-26, Figure 10a). 5 µl of each fraction was subjected to 15 % SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining. Active fractions eluted from heparin-Sepharose (Figure 10a) were pooled and concentrated (x 6) on YM3 cut-off membrane. 0.5 ml of the concentrated material was applied onto 30 ml Superdex 75 FPLC column equilibrated with 10 mM sodium acetate buffer, pH 5.0, containing 0.8 M NaCl, 1 mM DTT and 0.1 % CHAPS. Fractions (0.56 ml) were collected at a flow rate of 0.75 ml/min. Aliquots of each fraction were tested for heparanase activity and were subjected to SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining (Figure 11b).

## **EXAMPLE 1**

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#### Cloning of the hpa gene

[0124] Purified fraction of heparanase isolated from human hepatoma cells (SK-hep-1) was subjected to tryptic digestion and microsequencing. EST (Expressed Sequence Tag) databases were screened for homology to the back translated DNA sequences corresponding to the obtained peptides. Two EST sequences (accession Nos. N41349 and N45367) contained a DNA sequence encoding the peptide YGPDVGQPR (SEQ ID N0:8). These two sequences were derived from clones 257548 and 260138 (I.M.A.G.E Consortium) prepared from 8 to 9 weeks placenta cDNA library (Soares). Both clones which were found to be identical contained an insert of 1020 bp which included an open reading frame (ORF) of 973 bp followed by a 3' untranslated region of 27 bp and a Poly A tail. No translation start site (AUG) was identified at the 5' end of these clones.

[0125] Cloning of the missing 5' end was performed by PCR amplification of DNA from a placenta Marathon RACE (RIM) cDNA composite. A 900 bp fragment (designated hp3), partially overlapping with the identified 3' encoding EST clones was obtained

[0126] The joined cDNA fragment, 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes, as shown in Figure 1 and SEQ ID NO: 11, a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons. The 3' end of the partial cDNA inserts contained in clones 257548 and,260138 started, at nucleotide G<sup>721</sup> of SEQ ID NO:9 and Figure 1.

[0127] As further shown in Figure 1, there was a single sequence discrepancy between the EST clones and the PCR amplified sequence, which led to an amino acid substitution from Tyr<sup>246</sup> in the EST to Phe<sup>246</sup> in the amplified cDNA. The nucleotide sequence of the PCR amplified cDNA fragment was verified from two independent amplification products. The new gene was designated *hpa*.

[0128] As stated above, the 3' end of the partial cDNA inserts contained in EST clones 257548 and 260138 started at nucleotide 721 of hpa (SEQ ID NO:9). The ability of the hpa cDNA to form stable secondary structures, such as stem and loop structures involving nucleotide stretches in the vicinity of position 721 was investigated using computer modeling. It was found that stable stem and loop structures are likely to be formed involving nucleotides 698-724 (SEQ ID NO:9). In addition, a high GC content, up to 70 %, characterizes the 5' end region of the hpa gene, as compared to about only 40 % in the 3' region. These findings may explain the immature termination and therefore lack of 5' ends in the EST clones. [0129] To examine the ability of the hpa gene product to catalyze degradation of heparan sulfate in an in vitro assay the entire open reading frame was expressed in insect cells, using the Baculovirus expression system. Extracts of cells, infected with virus containing the hpa gene, demonstrated a high level of heparan sulfate degradation activity, while cells infected with a similar construct containing no hpa gene had no such activity, nor did non-infected cells. These results are further demonstrated in the following Examples.

## **EXAMPLE 2**

## Degradation of soluble ECM-derived HSPG

40 [0130] Monolayer cultures of High Five (RIM) cells were infected (72 h, 28 °C) with recombinant Bacoluvirus containing the pFasthpa plasmid or with control virus containing an insert free plasmid. The cells were harvested and lysed in heparanase reaction buffer by three cycles of freezing and thawing. The cell lysates were then incubated (18 h, 37 °C) with sulfate labeled, ECM-derived. HSPG (peak 1), followed by gel filtration analysis (Sepharose 6B) of the reaction mixture.

[0131] As shown in Figure 2, the substrate alone included almost entirely high molecular weight (Mr) material eluted next to  $V_0$  (peak I, fractions 5-20, Kav < 0.35). A similar elution pattern was obtained when the HSPG substrate was incubated with lysates of cells that were infected with control virus. In contrast, incubation of the HSPG substrate with lysates of cells infected with the *hpa* containing virus resulted in a complete conversion of the high Mr substrate into low Mr labeled degradation fragments (peak II, fractions 22-35, 0.5 < Kav < 0.75).

[0132] Fragments eluted in peak II were shown to be degradation products of heparan sulfate, as they were (i) 5- to 6-fold smaller than intact heparan sulfate side chains (Kav approx. 0.33) released from ECM by treatment with either alkaline borohydride or papain; and (ii) resistant to further digestion with papain or chondroitinase ABC, and susceptible to deamination by nitrous acid (6, 11).

[0133] Similar results (not shown) were obtained with Sf21 cells. Again, heparanase activity was detected in cells infected with the *hpa* containing virus (pF*hpa*), but not with control virus (pF). This result was obtained with two independently generated recombinant viruses. Lysates of control not infected High Five (RIM) cells failed to degrade the HSPG substrate.

[0134] In subsequent experiments, the labeled HSPG substrate was incubated with medium conditioned by infected

High Five (RIM) or Sf21 cells.

[0135] As shown in Figures 3a-b, heparanase activity, reflected by the conversion of the high Mr peak I substrate into the low Mr peak II which represents HS degradation fragments, was found in the culture medium of cells infected with the pFhpa2 or pFhpa4 viruses, but not with the control pF1 or pF2 viruses. No heparanase activity was detected in the culture medium of control non-infected High Five (RIM) or Sf21 cells.

[0136] The medium of cells infected with the pFhpa4 virus was passed through a 50 kDa cut off membrane to obtain a crude estimation of the molecular weight of the recombinant heparanase enzyme. As demonstrated in Figure 4, all the enzymatic activity was retained in the upper compartment and there was no activity in the flow through (<50 kDa) material. This result is consistent with the expected molecular weight of the hpa gene product.

[0137] In order to further characterize the *hpa* product the inhibitory effect of heparin, a potent inhibitor of heparanase mediated HS degradation (40) was examined.

[0138] As demonstrated in Figures 5a-b, conversion of the peak I substrate into peak II HS degradation fragments was completely abolished in the presence of heparin.

[0139] Altogether, these results indicate that the heparanase enzyme is expressed in an active form by insect cells infected with Baculovirus containing the newly identified human hpa. gene.

## **EXAMPLE 3**

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## Degradation of HSPG in intact ECM

[0140] Next, the ability of intact infected insect cells to degrade HS in intact, naturally produced ECM was investigated. For this purpose, High Five (RIM) or Sf21 cells were seeded on metabolically sulfate labeled ECM followed by infection (48 h, 28 °C) with either the pFhpa4 or control pF2 viruses. The pH of the medium was then adjusted to pH 6.2-6.4 and the cells further incubated with the labeled ECM for another 48 h at 28 °C or 24 h at 37 °C. Sulfate labeled material released into the incubation medium was analyzed by gel filtration on Sepharose 6B.

[0141] As shown in Figures- 6a-b and 7a-b, incubation of the ECM with cells infected with the control pF2 virus resulted in a constant release of labeled material that consisted almost entirely (>90%) of high Mr fragments (peak I) eluted with or next to Vo. It was previously shown that a proteolytic activity residing in the ECM itself and/or expressed by cells is responsible for release of the high Mr material (6). This nearly intact HSPG provides a soluble substrate for subsequent degradation by heparanase, as also indicated by the relatively large amount of peak I material accumulating when the heparanase enzyme is inhibited by heparin (6, 7, 12, Figure 9). On the other hand, incubation of the labeled ECM with cells infected with the pFhpa4 virus resulted in release of 60-70% of the ECM-associated radioactivity in the form of low Mr sulfate-labeled fragments (peak II, 0.5 <Kav < 0.75), regardless of whether the infected cells were incubated with the ECM at 28 °C or 37 °C. Control intact non-infected Sf21 or High Five (RIM) cells failed to degrade the ECM HS side chains. [0142] In subsequent experiments, as demonstrated in Figs. 8a-b, High Five (RIM) and Sf21 cells were infected (96 h, 28 °C) with pFhpa4 or control pF1 viruses and the culture medium incubated with sulfate-labeled ECM. Low Mr HS degradation fragments were released from the ECM only upon incubation with medium conditioned by pFhpa4 infected cells. As shown in Figure 9, production of these fragments was abolished in the presence of heparin. No heparanase activity was detected in the culture medium of control, non-infected cells. These results indicate that the lieparanase enzyme expressed by cells infected with the pFhpa4 virus is capable of degrading HS when complexed to other macromolecular constituents (i.e. fibronectin, laminin, collagen) of a naturally produced intact ECM, in a manner similar to that reported for highly metastatic tumor cells or activated cells of the immune system (6; 7).

## **EXAMPLE 4**

## Purification of recombinant heparanase

[0143] The recombinant heparanase was partially purified from medium of pFhpa4 infected Sf21 1 cells by Heparin-Sepharose chromatography (Figure 10a) followed by gel filtration of the pooled active fractions over an FPLC Superdex 75 column (Figure 11a). A ~ 63 kDa protein was observed, whose quantity, as was detected by silver stained SDS-polyacrylamide gel electrophoresis, correlated with heparanase activity in the relevant column fractions (Figures 10b and 11b, respectively). This protein was not detected in the culture medium of cells infected with the control pF1virus and was subjected to a similar fractionation on heparin-Sepharose (not shown).

## **EXAMPLE 5**

## Expression of the hpa gene in various cell types, organs and tissues

[0144] Referring now to Figures 12a-e, RT-PCR was applied to evaluate the expression of the *hpa* gene by various cell types and tissues. For this purpose, total RNA was reverse transcribed and amplified. The expected 585 bp long cDNA was clearly demonstrated in human kidney, placenta (8 and 11 weeks) and mole tissues, as well as in freshly isolated and short termed (1.5-48 h) cultured human placental cytotrophoblastic cells (Figure 12a), all known to express a high heparanase activity (41). The *hpa* transcript was also expressed by normal human neutrophils (Figure 12b). In contrast, there was no detectable expression of the *hpa* mRNA in embryonic human muscle tissue, thymus, heart and adrenal. (Figure 12b). The *hpa* gene was expressed by several, but not all, human bladder carcinoma cell lines (Figure 12c), SK hepatoma (SK-hep-1), ovarian carcinoma (OV 1063), breast carcinoma (435, 231), melanoma and megakary-ocytic (DAMI, CHRF) human cell lines (Figures 12d-e).

[0145] The above described expression pattern of the *hpa* transcript was determined to be in a very good correlation with heparanase activity levels determined in various tissues and cell types (not shown).

#### **EXAMPLE 6**

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#### hpa homologous genes

[0146] EST databases were screened for sequences homologous to the *hpa* gene. Three mouse ESTs were identified (accession No. Aa177901, from mouse spleen, Aa067997 from mouse skin, Aa47943 from mouse embryo), assembled into a 824 bp cDNA fragment which contains a partial open reading frame (lacking a 5' end) of 629 bp and a 3' untranslated region of 195 bp (SEQ ID NO:12). As shown in Figure 13, the coding region is 80% similar to the 3' end of the *hpa* cDNA sequence. These ESTs are probably cDNA fragments of the mouse *hpa* homolog that encodes for the mouse heparanase. [0147] Searching for consensus protein domains revealed an amino terminal homology between the heparanase and several precursor proteins such as Procollagen Alpha 1 precursor, Tyrosine-protein kinase-RYK, Fibulin-1, Insulin-like growth factor binding protein and several others. The amino terminus is highly hydrophobic and contains a potential trans-membrane domain. The homology to known signal peptide sequences suggests that it could function as a signal peptide for protein localization.

## **EXAMPLE 7**

## Isolation of an extended 5' end of hpa cDNA from human SK-hep1 cell line

[0148] The 5' end of hpa cDNA was isolated from human SK-hep1 cell line by PCR amplification using the Marathon RACE (RIM) (rapid amplification of cDNA ends) kit (Clontech). Total RNA was prepared from SK-hepl cells using the TRI-Reagent (Molecular research center Inc.) according to the manufacturer instructions. Poly A+ RNA was isolated using the mRNA separator kit (Clonetech).

[0149] The Marahton RACE SK-hep1 cDNA composite was constructed according to the manufacturer recommendations. First round of amplification was performed using an adaptor specific primer AP1: 5'-CCATCCTAATACG ACT-CACTATAGGGC-3', SEQ ID NO:1, and a hpa specific antisense primer hpl-629: 5'-CCCCAGGAGCAGCAGCAGCATCAG-3', SEQ ID NO:17, corresponding to nucleotides 119-99 of SEQ ID NO:9. The resulting PCR product was subjected to a second round of amplification using an adaptor specific nested primer AP2: 5'-ACTCACTATAGGGCTCGAGCGGC-3', SEQ ID NO:3, and a hpa specific antisense nested primer hpl-666 5'-AGGCTTCGAGCGCAGCAGCAT-3', SEQ ID NO:18, corresponding to nucleotides 83-63 of SEQ ID NO:9. The PCR program was as follows: a hot start of 94 °C for 1 minute, followed by 30 cycles of 90 °C - 30 seconds, 68 °C - 4 minutes. The resulting 300 bp DNA fragment was extracted from an agarose gel and cloned into the vector pGEM-T Easy (Promega). The resulting recombinant plasmid was designated pHPSK1.

[0150] The nucleotide sequence of the pHPSK1 insert was determined and it was found to contain 62 nucleotides of the 5' end of the placenta *hpa* cDNA (SEQ ID NO:9) and additional 178 nucleotides upstream, the first 178 nucleotides of SEQ ID NOs:13 and 15.

[0151] A single nucleotide discrepancy was identified between the SK-hepl 1 cDNA and the placenta cDNA. The "T" derivative at position 9 of the placenta cDNA (SEQ ID NO:9), is replaced by a "C" derivative at the corresponding position 187 of the SK-hep cDNA (SEQ ID NO: 13).

[0152] The discrepancy is likely to be due to a mutation at the 5' end of the placenta cDNA clone as confirmed by sequence analysis of sevsral additional cDNA clones isolated from placenta, which like the SK-hepl cDNA contained C at position 9 of SEQ ID NO:9..

[0153] The 5' extended sequence of the SK-hepl hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQIDNO:9). The assembled sequence contained an open reading frame which encodes, as shown in SEQ ID NOs:14 and 15, a polypeptide of 592 amino acids with a calculated molecular weight of 66,407 daltons. The open reading frame is flanked by 93 bp 5' untranslated region (UTR).

#### **EXAMPLE 8**

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## Isolation of the upstream genomic region of the hpa gene

[0154] The upstream region of the *hpa* gene was isolated using the Genome Walker kit (Clontech) according to the manufacturer recommendations. The kit includes five human genomic DNA samples each digested with a different restriction endonuclease creating blunt ends: *EcoRV*, *Scal*, *Dral*, *Pvull* and *Sspl*.

[0155] The blunt ended DNA fragments are ligated to partially single stranded adaptors. The Genomic DNA samples were subjected to PCR amplification using the adaptor specific primer and a gene specific primer. Amplification was performed with Expand High Fidelity (Boehringer Mannheim).

[0156] A first round of amplification was performed using the ap1 primer: 5'-G TAATACGACTCACTATAGGGC-3', SEQ ID NO:19, and the *hpa* specific antisense primer hpl-666: 5'-AGGCTTCGAGCGCAGCAGCAT-3', SEQ ID NO:18, corresponding to nucleotides 83—63 of SEQ ID NO:9. The PCR program was as follows: a hot start of 94 °C - 3 minutes, followed by 36 cycles of 94 °C - 40 seconds, 67 °C - 4 minutes.

[0157] The PCR products of the first amplification were diluted 1:50. One µI of the diluted sample was used as a template for a second amplification using a nested adaptor specific primer ap2: 5'-ACTATAGGGCACGCGTGGT-3', SEQ ID NO:20, and a hpa specific antisense primer hpI-690, 5'-CTTGGGCTCACC TGGCTGCTC-3', SEQ ID NO:21, corresponding to nucleotides 62-42 of SEQ ID NO:9. The resulting amplification products were analyzed using agarose gel electrophoresis. Five different PCR products were obtained from the five amplification reactions. A DNA fragment of approximately 750 bp which was obtained from the Sspl digested DNA sample was gel extracted. The purified fragment was ligated into the plasmid vector pGEM-T Easy (Promega). The resulting recombinant plasmid was designated pGHP6905 and the nucleotide sequence of the hpa insert was determined.

[0158] A partial sequence of 594 nucleotides is shown in SEQ ID NO:16. The last nucleotide in SEQ ID NO:13 corresponds to nucleotide 93 in SEQ ID:13. The DNA sequence in SEQ ID NO:16 contains the 5' region of the *hpa* cDNA and 501 nucleotides of the genomic upstream region which are predicted to contain the promoter region of the *hpa* gene.

## **EXAMPLE 9**

## 35 Expression of the 592 amino acids HPA polypeptide in a human 293 cell line

[0159] The 592 amino acids open reading frame (SEQ ID NOs:13 and 15) was constructed by ligation of the 110 bp corresponding to the 5' end of the SK-hep 1 hpa cDNA with the placenta cDNA. More specifically the Marathon RACE (RIM) PCR amplification product of the placenta hpa DNA was digested with SacI and an approximately 1 kb fragment was ligated into a SacI-digested pGHP6905 plasmid. The resulting plasmid was digested with Earl and AatII. The Earl sticky ends were blunted and an approximately 280 bp Earl/blunt-AatII fragment was isolated. This fragment was ligated with pFasthpa digested with EcoRI which was blunt ended using Klenow fragment and further digested with AatII. The resulting plasmid contained a 1827 bp insert which includes an open reading frame of 1776 bp, 31 bp of 3' UTR and 21 bp of 5' UTR. This plasmid was designated pFastLhpa.

[0160] A mammalian expression vector was constructed to drive the expression of the 592 amino acids heparanase polypeptide in human cells. The hpa cDNA was excised prom pFastLhpa with BssHII and Noti. The resulting 1850 bp BssHII-Noti fragment was ligated to a. mammalian expression vector pSI (Promega) digested with Mlul and Noti. The resulting recombinant plasmid, pSIhpaMet2 was transfected into a human 293 embryonic kidney cell line.

[0161] Transient expression of the 592 amino-acids heparanase was examined by western blot analysis and the enzymatic, activity was tested using the gel shift assay. Both these procedures are described in length in U.S. Pat. application No. 09/071,739, filed May 1, 1998, which is incorporated by reference as if fully set forth herein. Cells were harvested 3 days following transfection. Harvested cells were re-suspended in lysis buffer containing 150 mM NaCl, 50, mM Tris pH 7.5, 1% Triton X-100, 1 mM PMSF and protease inhibitor cocktail (Boehringer Mannheim). 40 μg protein extract samples were used for separation on a SDS-PAGE. Proteins were transferred onto a PVDF Hybond-P membrane (Amersham). The membrane was incubated with an affinity purified polyclonal anti heparanase antibody, as described in U.S. Pat. application No. 09/071,739. A major band of approximately 50 kDa was observed in the transfected cells as well as a minor band of approximately 65 kDa. A similar pattern was observed in extracts of cells transfected with the pShpa as demonstrated in U.S. Pat application No. 09/071,739. These two bands probably represent two forms of

the recombinant heparanase protein produced by the transfected cells. The 65 kDa protein probably represents a heparanase precursor, while the 50 kDa protein is suggested herein to be the processed or mature form.

[0162] The catalytic activity of the recombinant protein expressed in the pShpaMet2 transfected cells was tested by gel shift assay. Cell extracts of transfected and of mock transfected cells were incubated overnight with heparin (6 µg in each reaction) at 37 °C, in the presence of 20 mM phosphate citrate buffer pH 5.4, 1 mM CaCl<sub>2</sub>, 1 mM DTT and 50 mM NaCl. Reaction mixtures were then separated on a 10 % polyacrylamide gel. The catalytic activity of the recombinant heparanase was clearly demonstrated by a faster migration of the heparin molecules incubated with the transfected cell extract as compared to the control. Faster migration indicates the disappearance of high molecular weight heparin molecules and the generation of low molecular weight degradation products.

#### **EXAMPLE 10**

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#### Chromosomal localization of the hpa gene

15 [0163] Chromosomal mapping of the *hpa* gene was performed utilizing a panel of monochromosomal human/CHO and human/mouse somatic cell hybrids, obtained from the UK HGMP Resource Center (Cambridge, England).

[0164] 40 ng of each of the somatic cell hybrid DNA samples were subjected to PCR amplification using the *hpa* primers: hpu565 5'-AGCTCTGTAGATGTGC TATACAC-3', SEQ ID NO:22, corresponding to nucleotides 564-586 of SEQ ID NO:9 and an antisense primer hpl171 5'-GCATCTTAGCCGTCTTTCTTCG-3', SEQ ID NO:23, corresponding to nucleotides 897-876 of SEQ ID NO:9.

[0165] The PCR program was as follows: a hot start of 94 °C - 3 minutes, - followed by 7 cycles of 94 °C - 45 seconds, 66 °C - 1 minute, 68 °C - 5 minutes, followed by 30 cycles of 94 °C - 45 seconds, 62 °C - 1 minute, 68 °C - 5 minutes, and a 10 minutes final extension at 72 °C.

[0166] The reactions were performed with Expand long PCR (Boehringer Mannheim). The resulting amplification products were analyzed using agarose gel electrophoresis. As demonstrated in Figure 14, a single band of approximately 2.8 Kb was obtained from chromosome 4, as well as from the control human genomic DNA. A 2.8 kb amplification product is expected based on amplification of the genomic *hpa* clone (data not shown). No amplification products were obtained neither in the control DNA samples of hamster and mouse nor in somatic hybrids of other human chromosome.

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#### SEQUENCE LISTING

[0168]

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- (1) GENERAL INFORMATION:
- (i) APPLICANT: Iris Pecker, Israel VLodavsky and Elena Feinstein
  - (ii) TITLE OF INVENTION: POLYNUCLEOTIDE ENCODING A POLYPEPTIDE HAVING HEPARANASE ACTIVITY AND EXPRESSION OF SAME IN TRANSDUCED CELLS
  - (iii) NUMBER OF SEQUENCES: 23
  - (iv) CORRESPONDENCE ADDRESS:

- (A) ADDRESSEE: Mark M. Friedman c/o Robert Sheinbein
- (B) STREET: 2940 Birchtree lane
- (C) CITY: Silver Spring (D) STATE: Maryland
- (E) COUNTRY: United States of America
- (F) ZIP: 20906
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: 1.44 megabyte, 3.5" microdisk
  - (8) COMPUTER: Twinhead\* Slimnote-890TX
  - (C) OPERATING SYSTEM: MS DOS version 6.2, Windows version 3.11
  - (D) SOFTWARE: Word for Windows version 2.0 converted to an ASCI file
- 35 (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER:
  - (B) FILING DATE:
  - (C) CLASSIFICATION:
  - (vii) PRIOR APPLICATION DATA:
    - (A) APPLICATION NUMBER: 08/922,170
    - (B) FILING DATE: 2 SEP 1997
  - (viii) ATTORNEY/AGENT INFORMATION:
    - (A) NAME: Friedmam, Mark M.
    - (B) REGISTRATION NUMBER: 33,883
  - (C) REFERENCE/DOCKET NUMBER: 910/1
    - (ix) TELECOMMUNICATION INFORMATION:
      - (A) TELEPHONE: 972-3-5625553
      - (B) TELEFAX: 972-3-5625554
      - (C) TELEX:
  - (2) INFORMATION FOR SEQ ID N0:1:

5	(A) LENGTH: 27 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D.) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEC	Q ID N0:1:
10	c	CATCCTAAT ACGACTCACT ATAGGGC 2
	(2) INFORMATION FOR SEQ ID NO:2:	
15	(i) SEQUENCE CHARACTERISTICS	:
20	(A) LENGTH: 24 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEC	Q ID NO:2:
25		GTAGTGATGC CATGTAACTG AATC 24
20	(2) INFORMATION FOR SEQ ID NO:3:	
	(i) SEQUENCE CHARACTERISTICS	:
30	(A) LENGTH: 23 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	·
35	(xi) SEQUENCE DESCRIPTION: SEC	D ID NO:3:
		ACTCACTATA GGGCTCGAGC GGC 23
40		ACTUACIATA GUGETEGAGE GGE 23
	(2) INFORMATION FOR SEQ ID NO:4:	
	(i) SEQUENCE CHARACTERISTICS	
45	<ul><li>(A) LENGTH: 22</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
50	(xi) SEQUENCE DESCRIPTION: SEC	Q ID NO:4:
		GCATCTTAGC CGTCTTTCTT CG 22
55	(2) INFORMATION FOR SEO ID NO:5:	
	(i) SEQUENCE CHARACTERISTICS:	

(i) SEQUENCE CHARACTERISTICS:

	(A) LENGTH: 15 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
5	(xi) SEQUENCE DESCRIPTION: SE	EQ 10 NO:5:
10	(2) INFORMATION FOR SEQ 10 NO:6:	
	(i) SEQUENCE CHARACTERISTICS	<b>S</b> :
15	<ul><li>(A) LENGTH: 23</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
20	(xi) SEQUENCE DESCRIPTION: SE	Q ID NO:6:
		TTCGATCCCA AGAAGGAATC AAC 23
25	(2) INFORMATION FOR SEQ ID NO:7:	
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40	(2) INFORMATION FOR SEQ 10 NO:8:	
40	(i) SEQUENCE CHARACTERISTICS	<b>S</b> :
45	<ul><li>(A) LENGTH: 9</li><li>(B) TYPE: amino acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
	(xi) SEQUENCE DESCRIPTION: SE	Q ID N0:8:
50	ту	r Gly Pro Asp Val Gly Gln Pro Arg 5
	(2) INFORMATION FOR SEO ID NO:9:	
55	(i) SEQUENCE CHARACTERISTICS	S:
	(A) LENGTH: 1721 (B) TYPE: nucleic acid	

(C) STRANDEDNESS: double (D) TOPOLOGY: linear

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## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

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TCAAGAACAG CACCTACTCA AGAAGCTCTG TAGATGTGCT ATACACTTTT GCAAACTGCT 600 CAGGACTGGA CTTGATCTTT GGCCTAAATG CGTTATTAAG AACAGCAGAT TTGCAGTGGA 660 ACAGTICTAA IGCICAGTIG CICCIGGACT ACTGCTCTIC CAAGGGGTAT AACATTICTI 720 GGGAACTAGG CAATGAACCT AACAGTTICC TTAAGAAGGC TGATATTITC ATCAATGGGT 780 CGCAGTTAGG AGAAGATTAT ATTCAATTGC ATAAACTTCT AAGAAAGTCC ACCTTCAAAA 840 ATGCAAAACT CTATGGTCCT GATGTTGGTC AGCCTCGAAG AAAGACGGCT AAGATGCTGA 900 AGAGCTICCI GAAGGCTGGT GGAGAAGTGA TIGATTCAGT TACATGGCAT CACTACTATT 960 TGAATGGACG GACTGCTACC AGGGAAGATT TTCTAAACCC TGATGTATTG GACATTITTA 1020 TITCATCTGT GCAAAAAGTT TTCCAGGTGG TTGAGAGCAC CAGGCCTGGC AAGAAGGTCT 1080 GGTTAGGAGA AACAAGCTCT GCATATGGAG GCGGAGCGCC CTTGCTATCC GACACCTTTG 1140 CAGCTGGCTT TATGTGGCTG GATAAATTGG GCCTGTCAGC CCGAATGGGA ATAGAAGTGG 1200 TGATGAGGCA AGTATTCTTT GGAGCAGGAA ACTACCATTT AGTGGATGAA AACTTCGATC 1260 CTITACCIGA TTATIGGCIA TCICTICIGI TCAAGAAATI GGTGGGCACC AAGGTGTIAA 1320 TGGCAAGCGI GCAAGGTICA AAGAGAAGGA AGCTYCGAGI ATACCTICAT IGCACAAACA 1380 CTGACAATCC AAGGTATAAA GAAGGAGATT TAACTCTGTA TGCCATAAAC CTCCATAACG 1440 TCACCAAGTA CTTGCGGTTA CCCTATCCTT TTTCTAACAA GCAAGTGGAT AAATACCTTC 1500 TAAGACCTTT GGGACCTCAT GGATTACTTT CCAAATCTGT CCAACTCAAT GGTCTAACTC 1560 TAAAGATGGT GGATGATCAA ACCTTGCCAC CTTTAATGGA AAAACCTCTC CGGCCAGGAA 1620 GIICACIGGG CIIGCCAGCI TICYCATATA GIITIIITGI GAIAAGAAAI GCCAAAGIIG 1680 CIGCTIGCAT CIGAAAATAA AATATACTAG TCCTGACACT G

(2) INFORMATION FOR SEQ ID NO:10:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 543

(B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ 10 NO:10:

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5	l eu	Leu	Gly	Pro 20		Gly	Pro	Leu	Ser 25	Рго	Gly	Ala	Leu	Pro 30	_	Pro
	Ala	Gln	Ala 35		Asp	Val	Val	Asp 40		Asp	Phe	Phe	Thr 45		Glu	Pro
	Leu	His 50		Val	Ser	Pro	Ser 55	Phe	Leu	Ser	Val	Thr 60	He	Asp	Ala	Asn
10	Leu 65			Asp	Pro	Arg 70	Phe	Leu	He	Leu	Leu 75		Ser	Pro	Lys	Leu 80
		Thr	Leu	Ala				Ser	Pro			Leu	Arg	Phe		
15	Thr	Lys	Thr		B5 Phe	Leu	lle	Phe		90 Pro	lys	Lys	Glu	Ser	95 Thr	Phe
	Glu	Glu	Arg	100 Ser	Туг	Τrp	Gln	Ser	105 Gln	Val	Asn	Gln	Asp	110 11e	Cys	Lys
20	Tyr	Gly	115 Ser	ile	Pro	Pro	Asp	120 Val	Glu	Glu	Lys	Leu	125 Arg	Leu	Glu	Trp
		130					135	Leu				140				
25	145					150		Ser			155					160
25					165					170					175	
				180				Leu	185					190		
30	Arg	Thr	Al 8 195	Asp	Leu	Gln	тгр	Asn 200	Ser	Ser	Asn	Ala	Gln 205	Leu	Leu	Leu
. •	•	Tyr 210	Cys	Ser	Ser	Lys	Gly 215	Tyr	Asn	Ile	Ser	7 rp 220	FLU	Leu	GLY	Asn
35	Glu 225	Pro	Asn	Ser	Phe	Leu 230	Lys	Lys	Ala	Asp	11e 235	Phe	lle	Asn	Gly	Ser 240
	Gln	Leu	Gly	Głu	Asp 245	Tyr	ile	Gln	Leu	His 250	Lys	Leu	Leu	Arg	Lys 255	Ser
40	Thr	Phe	Lys	Asn 260	Ala	Lys	Leu	Туг	265 265	Pro	Asp	Val	Gly	Gln 270	Pro	Arg
	Arg	•	1hr 275	Ala	Lys	Met	Leu	280 280	\$er	Phe	Leu		Ala 285	Gly	Gly	Glu
	Val	11e 290	Asp	Ser	Val	Thr	1rp 295	His	His	ТУГ		Leu 300	Asn	Gly	Arg	Thr
45	Ala 305	Thr	Arg	Glu		Phe 310	Leu	Asn	Рго	Asp	Val 315	Leu	Asp	ile	Phe	1 l e 320
	Ser	Ser	Val		Lys 325	Val	Phe	Gln		Va l 330	Glu	Ser	Thr	Arg	Pro 335	Gly
50	Lys	Lys		1rp 340	f en	Gly	Glu		Ser 345	Ser	Ala	Tyr	Gly	Gly 350	Gly	Ala
	Pro				Asp	Thr	Phe	ala 088	Ala	Gly	Phe	Het	1rp 365		Asp	Lys
55	Leu			Ser	Ala	Arg	He t		lle	Glu	Val	Val		Arg	Gln	Val

			370					375					380				
5		Phe 385	Phe	Gly	Ala	Gly	Asn 390	Tyr	His	Leu	val	Asp 395	Glu	Asn	Phe	Asp	Pro 400
5		Leu	Pro	Asp	Туг	Trp 405	Leu	Ser	Leu	Leu	Phe 410	Lys	Lys	Leu	Val	Gly 415	Thr
		lys	Val	Leu	Met 42	Ala D	Ser	Val	Gln	Gly 425	Ser 5	Lys	Arg	Arg	Lys 430	L eu )	Arg
10		Val	Туг	Leu 435	Hís	Cys	Thr	Asn	Thr 440	Asp	Asn	Pro	Arg	Tyr 445	Lys	Glu	Gly
		Asp	Leu 450	Thr	Fen	Tyr	Ala	11e 455	Asn	Leu	His	Asn	Val 460	Thr	Lys	Tyr	Leu
15		Arg 465	Leu	Pro	Tyr	Pro	Phe 470	Ser	Asn	Lys	Gln	Val 475	Asp	Lys	Туг	Leu	L eu 480
		Arg	Pro	Leu	Gly	Pro 485	His	Gly	Leu	Leu	Ser 490	Lys	Ser	Val	Gln	Leu 495	Asn
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		Glu	Lys	Pro 515	Leu	Arg	Рго	Gly	Ser 520	Ser	Leu	Gly	Leu	Рго 525	Ala	Phe	Ser
25		Tyr	Ser 530	Phe	Phe	Val	1 le	Arg 535	Asn	Ala	Lys	Val	Ala 540	Ala	Cys	1 l e 543	
	(2) INFORMATION FO	R SI	EO II	D NC	D:11:												
	, ,																
30	(i) SEQUENCE CF	IAR	ACTI	ERIS	TICS	S:											
	(A) LENGTH: (B) TYPE: nuc																
	(C) STRAHDE	DHE	ESS:		ble												
35	(D) TOPOLOG	iY: li	near	•													
	(xi) SEQUENCE D	ESC	RIP	TION	N: SE	Q ID	NO:	11:									

								•	•			CT	AGA	GCT	TTC	GAC	14
	TCT	CCG	CTG	CGC	GGC	AGC	TGG	CGG	GGG	GAG	CAG	CCA	GGT	GAG	ccc	AAG	62
5	ÀTG Met	ctc	ara	ćec"	TCG.	AAG	CCT	GCG	CTG	CCG	CCG	CCG	CTG	ATG	CTG	CTG	110
10	Leu	Leu	Gly	CCG Pro 20	Leu	Gly	Pro	Leu	Ser 25	Pro	Gly	Ala	Leu	Pro 30	Arg	Pro	158
	Ala	Gln	A1a 35	CAG Gln	Asp	Val	Val	Asp 40	Leu	Asp	Phe	Phe	1hr 45	Gln	Glu	Pro	206
15	Leu	His 50	Leu	GTG Val	Ser	Рго	Ser 55	Phe	Leu	Ser	Val	Thr 60	He	Asp	Ala	Asn	
20	Leu 65	Ala	Thr	GAC Asp	Pro	Arg 70	Phe	Leu	lle	Leu	75	Gly	Ser	Pro	Lys	80	302
	Arg	Thr	Leu	GCC	Arg 85	Gly	Leu	Ser	Pro	Ala 90	Tyr	Lev	Arg	Phe	Gly 95	Gly	350
25	Thr	Lys	Thr	100	Phe	Leu	ile	Phe	Asp 105	Pro	Lys	Lys	Glu	5er 110	Thr		398
	GAA Glu	GAG Głu	AGA Arg 115	Ser	TAC	TGG Trp	Gla	TCT Ser 120	Gln	GTC	ASO	Gln	GAT Asp 125	ile	TGC Cys	AAA Lys	446
30																	
35																	
40																	
45																	
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			Ser					Val		GAG Glu			Arg				494
5		Tyr			Gln		Leu			GAA Glu							
10										GTA Val 170							
										TTT Phe							638
15	Arg	Thr	Ala 195	Asp	Leu	Gln	Тгр	Asn 200	Ser	Ser	Asn	Ala	Gln 205	Leu	Leu	Leu	
	Asp	Tyr 210	Cys	Ser	Ser	Lys	Gly 215	ТУГ	Asn	att	\$er	1rp 220	Glu	Leu	Gly	Asn	734
20	Glu 225	Pro	Asņ	Ser	Phe	Leu 230	Lys	Lys	Ala	GAT ASP	11e 235	Phe	He	Asn	Gly	Ser 240	
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	Thr ; :	Phe	Lys	Asn 260	Ala	Lys	Leu	Tyr	Gly 265	Pro	Asp	Val	Gly	Gln 270	Pro	Arg	878 926
30	Arg <sup>*</sup>	Lys	1hr 275	Ala	Lys	Het	Leu	Lys 280	Ser	Phe	Leu	lys	Ala 285	Gly	Gly	Glu	974
	Val	1 l e 290	Asp	Ser	Val	Thr	7rp 295	His 	His -	Туг	Туг	Leu 300	Asn	Gly	Arg	Thr	1019
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40	Ser	Ser	Val	Gln	Lys 325	Val	Phe	Gln	Val	Val 330	Glu	Ser	Thr	Arg	Pro <b>33</b> 5	Gly	
	Lys			340					345					350			1163
45	Pro.	GGC	355 CTG	TCA I	GCC	CGA	ATG	360 GGA	ATA	GAA	GTG	GTG	365 ATG	AGG	CAA	GTA	1211
	TTC	370 111 -	GGA I	GCA (	GGA	AAC	375 TAC	CAT	TTA	GTG	GAT	380 GAA	AAC	TTC	GAT	133	1259
50		CCT :	GAT	TAT	TGG	390 CTA	TCT	CTT	CTG	TTC	395 AAG	AAA	TTG	GTG	GGC	400 ACC	1307
55	AAG	GTG	TTA :	ATG (	405 GCA	AGC	GTG	CAA	GGT	410 TCA	AAG	AGA	AGG	AAG	415 CTT	CGA	1355
	Lys	Val	l eu	Het	6 I A	ser	۷ə۱	Gln	Gly	Ser	lys	Arg	Arg	L ys	reu	Arg	

					420					425					430			
5																GAA Glu		1403
		Asp																1451
10		CGG Arg 465																1499
15		AGA   Arg																1547
		GGT (																1595
20		GAA /	Lys															1643
25		TAT /															TGA	
23		AAA 1	TAA	AAT	ATA	CTA	GTC	CTG	ACA	CTG								1718
	(2) INFORMATION	FOR -	SEC	) ID I	NO:1	12:												
30	(i) SEQUENCE	CHAI	RAC	TER	RISTI	CS:												
35	(A) LENGT (B) TYPE: (C) STRAN (D) TOPOL	nuclei IDEDI	c ac	S: do	ouble	e												
	(xi) SEQUENCI	E DES	SCR	IPTIO	ON:	SEQ	ID N	O:12	2									
40	1	TGGCA IGTCCA IGGGCA ATGAAA GTCCCA	TAG	C CT A AG	TTGC TCGT AGCC	AGCT GATG TTTA	GGC AGG CCT	TTTA CAGG GATT	TGT TGT ACT	GGCT TCTT GGCT	GGAT CGGA CTCT	AA A GC A CT T	TTGG GGCA CTGT	GCCT ACTA TCAA	G TC C CA G AA	AGCC CTTA ACTG	CAGA GTGG GTAG	120 180 240
45	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	CCACT GAACC GGATA GAACG CTCTCC GAAATG	GCAI CGT/ GTC/ CCGI CCGI	C TA A TA A CC A AA C AG A AA G GG	ACGT ATGT TTCT TTCT GAAG TCGC GTGT	CTAT CACC GAAG GAAG TGCA TGCT TATT	CAC AAG CCT ATG CTA TGT	CCAC CACT TCGG GTGG AGCC ATAT AAAA	GAT TGA GGC ATG TGC GAA	ATCA AGGT CGGA AGCA CTGC AATA AACC	GGAA ACCG TGGA GACC CTTT AAAG CTAG	GG A CC T TT A CT G TC C GC A TT T	GATC CCGT CTTT CCAG TATG TACG AGGA	TAAC TGTT CCAA CTTT GTTT GTAC GGCC	T CT C AG A TC G AC T TT C CC	GTAT GAAA TGTC AGAA TGTC TGAG	GTCC CCAG CAAC AAAC ATAA ACAA TGCC	360 420 480 540 600 660 720
50		AGTTO												ICAG	1 61	6616	1161	824
	(2) INFORMATION	FOR:	SEQ	l D I	NO:1	3:												
	(i) SEQUENCE	CHA	RAC	TER	ISTI	CS:												
55	(A) LENGT (B) TYPE: r (C) STRAN	nuclei	c aci		ouble	<b>:</b>												
	·																	

## (D) TOPOLOGY: linear

# (xi) SEQUENCE DESCRIPTION: SEO ID NO:13

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	MIGGIOGO	300
10	C1001000 101000100 001001000 001001000 1010001000	360
	Cidatelie icateman appearance cidalation entre continue	420
	Wilderson were an expense of the contract of t	480
	COLUCTION COMPANY AND	540
	I I CONTROLLE CONTROL CONTROL NOT THE PROPERTY OF THE PROPERTY	600
	CANDICANCE NOBLINITION TO CONTROL OF THE CONTROL OF	660
15	CGGTIGGAAT GGCCCTACCA GGAGCAATIG CTACTCCGAG AACACTACCA GAAAAAGTIC	720
15		
	AAGAACAGCA CCTACTCAAG AAGCTCTGTA GATGTGCTAT ACACTTTTGC AAACTGCTCA	780
	GGACTGGACT TGATCTTIGG CCTAAATGCG TTATTAAGAA CAGCAGATTI GCAGTGGAAC	840
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		1200
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	TTAGGAGAAA CAAGCTCTGC ATATGGAGGC GGAGCGCCCT TGCTATCCGA CACCTTTGCA	1320
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	MUNCOTTUD CHOOLOMICS MITTER TO THE TELEVISION OF	1800
		1860
		1899
	de l'addition de l'addition de la company	

## (2) INFORMATION FOR SEQ ID NO:14:

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# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 592 (B) TYPE: amino acid (C) STRANDEDNESS: singl (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14

Het Glu Gly Ala Val Gly Gly Val Arg Arg Arg Asn Gly Ala Glu Glu Arg Arg Lys Gly Arg Trp Gly Ser Ala Gly Gly Ser Ala Arg 20 25 30 Ala Leu Asp Ser Pro Leu Arg Gly Ser Trp Arg Gly Glu Gln Pro Gly Glu Pro Lys Met Leu Leu Arg Ser Lys Pro Ala Leu Pro Pro : 50 Pro Leu Met Leu Leu Leu Gly Pro Leu Gly Pro Leu Ser Pro . 65 Gly Ala Leu Pro Arg Pro Ala Gln Ala Gln Asp Val Val Asp Leu Asp Phe Phe Thr Gln Glu Pro Leu His Leu Val Ser Pro Ser Phe Leu Ser Val Thr Ile Asp Ala Asn Leu Ala Thr Asp Pro Arg Phe .. 115. Leu lle Leu Leu Gly Ser Pro Lys Leu Arg Thr Leu Ala Arg Gly
125 130 Leu Ser Pro Ala Tyr Leu Arg Phe Gly Gly Thr Lys Thr Asp Phe Leu Ile Phe Asp Pro Lys Lys Glu Ser Thr Phe Glu Glu Arg Ser 155 160 165 Tyr Trp Gln Ser Gln Val Asn Gln Asp Ile Cys Lys Tyr Gly Ser Ile Pro Pro Asp Val Glu Glu Lys Leu Arg Leu Glu Trp Pro Tyr Gln Glu Gln Leu Leu Leu Arg Glu His Tyr Gln Lys Lys Phe tys Asn Ser Thr Tyr Ser Arg Ser Ser Val Asp Val Leu Tyr Thr Phe Ala Asn Cys Ser Gly Leu Asp Leu Ile Phe Gly Leu Asn Ala Leu Leu Arg Thr Ala Asp Leu Gln Trp Asn Ser Ser Asn Ala Gln Leu Leu Leu Asp Tyr Cys Ser Ser Lys Gly Tyr Asn Ile Ser Trp Glu Leu Gly Asn Glu Pro Asn Ser Phe Leu Lys Lys Ala Asp 1le Phe 275 280 285 He Ash Cly Ser Cln Leu Gly Glu Asp Tyr He Gin Leu His Lys
290 205 300 Leu Leu Arg Lys Ser Thr Phe Lys Asn Ala Lys Leu Tyr Gly Pro Asp Val Gly Gln Pro Arg Arg Lys Thr Ala Lys Met Leu Lys Ser Phe Leu Lys Ala Gly Gly Glu Val Ile Asp Ser Val Thr Trp His

				350					355		Glu			360
Asn	Pro	Asp	Val	Leu 365	Asp	He	Phe	He	Ser 370	Ser	Val	Gln	Lys	Val 375
				380					385		Lys			390
Gly	Glu	Thr	Ser	Ser 395	Ala	Tyr	Gly	Gly	Gly 400	Ala	Pro	Leu	Leu	Ser 405
Asp	Thr	Phe	Ala	Ala 410	Gly	Phe	Met	Тгр	Leu 415	Asp	Lys	Leu	Gly	Leu 420
				Gly 425					430		Gln			435
				440					445		Phe			450
Pro	Asp	Tyr	Trp	Leu 455	Ser	Leu	Leu	Phe	Lys 460	Lys	Leu	Val	Gly	Thr 465
				470					475		Arg			480
		•		485					490		Pro			495
		-		500					505		His			510
•	-		-	515			•		520		Lys			525
				530					535		Leu			540
				545					550		Val			555
				560					565		Pro			570
				575			Tyr	Ser	Phe 580	Phe	Val	Ile	Arg	Asn 585
Ala	Lys	Val	Ala	Ala 590	Cys	1 le 592								

(2) INFORMATION FOR SEO ID NO:15:

- (i) SEQUENCE CHARACTERISTICS:

  - (A) LENGTH: 1899 (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15

				. :		•									GGG	3
(A)		ccc	ACC:	AAG	CAA	GYA	CCA	GAG	AGC	CGG	GCA	GGC	999	GCG	GGG	48
	7.7.C	CAT	TCC	CAC	CAG	TGG	GAG	GGA	TGC	AGA	AGA	GGA	GTG	GGA	666	93
λ.	110	CAI	566	CCA	CTC	CCA	CCC	CTG	AGG	ACC	CGT	AAC	ccc	ece	GAG	138
	AIG	LAG.	CL	AL.	Wet	CL	CL	Val	Ara	Ara	Arg	Acn	CIV	Ala	Glu	,,,,
	met	Glu	GIY	ALB	5	uly	GLY	vat	nı y	10	vi A	7311	uty	~10	15	
4	GAA	AGG	AGA	AAA	GGG	CGC	TGG	GGC	TCG	GCG	GGA	GGA	AGT	GCT	AGA	183
	Glu	Arg	Arg	Lys	Gly	Arg	Trp	Gly	Ser	Ala	Gly	Gly	Ser	Ala	Arg	
			_	·	50					25					30	
	CCT	ctc	GAC	TCT	ccg	CTG	CGC	GGC	AGC	TGG	CGG	GGG	GAG	CAG	CCA	228
	ΔIs	Leu	Asp	Ser	Pro	Leu	Arg	GLY	Ser	Trp	Arg	Gly	Glu	Gln	Pro	
•			ЛОР		35		3	• • •		40	_	·			45	
	GGT	GAG	CCC	AAG	ATG	CTG	CTG	CGC	TCG	AAG	CCT	GCG	CTG	CCG	CCG	273
	Gly	Glu	Pro	Lys	Het	Leu	Leu	Arg	Ser	Lys	Pro	Ala	Leu	Pro	Pro	
					50					55				-	60	
	ccs	CTG	ATG	CTG	CTG	CTC	CTG	GGG	CCG	CTG	GGT	CCC	CTc	TCC	CCT	318
\i	Pro	Leu	Met	Leu	Leu	Leu	Leu	Gly	Pro	Leu	Gly	Рго	Leu	Ser	Pro	
					65			-		70					75	
	ccr	ccc	CIG	CCC	CGA	CCT	GCG	CAA	GCA	CAG	GAC	GTc	GTG	GAC	CTG	363
	CLV	415	Leu	Pro	Ara	Pro	Ala	Gln	Ala	Gln	Asp	Val	Vat	ASD	Leu	
	J.,	~ 10			80			_		85	•			·	90	
	GAC	TIC	TTC	ACC	CAG	GAG	CCG	CTG	CAC	CIG	GTG	AGC	CCC	TCG	TTC	408
	Asp	Phe	Phe	Thr	Gln	Glu	Pro	Leu	His	Leu	Val	Ser	Pro	Ser	Phe	
	,				95					100					105	
	CIG	TCC	GIC	ACC	ATT	GAC	GCC	AAC	CTG	GCC	ACG	GAC	CCG	CGG	TTC	453
						_										

	l eu	Ser	yal	Thr	11e 110	Asp	Ala	Asn	Lev	Ala 115	Thr	Asp	Pro	Arg	Phe 120	
5	r en C1C	ATC	CTC Leu	CTG Leu	GGT Gly 125	TCT Ser	CCA Pro	AAG Lys	CTT Leu	CGT Arg 130	ACC Thr	TTG Leu	GCC Ala	AGA Arg	GGC Gly 135	498
	I TG Leu	1CT Ser	CCT Pro	GCG Ala	TAC Tyr 140	Len C1è	AGG Arg	TTT Phe	GGT Gly	GGC Gly 145	ACC Thr	AAG Lys	ACA Thr	GAC Asp	TTC Phe 150	543
10	CTA Leu	ATT Ile	TTC Phe	GAT ASP	CCC Pro 155	AAG Lys	AAG Lys	GAA Glu	TCA Ser	ACC Thr 160	TTT Phe	GAA Glu	GAG Glu	AGA Arg	AGT Ser 165	588
15	Туг	Trp	Gln	TCT Ser	Gln 170	Val	Asn	Gln	Asp	11e 175	Cys	Lys	Tyr	Gly	Ser 180	633
	He	Pro	Pro	GAT Asp	Val 185	Glu	Glu	Lys	Leu	Arg 190	Leu	Glu	Trp	Pro	Туг 195	678
20	Gln	Glu	Gln	TTG Leu	Leu 200	Leu	Arg	Glu	His	7yr 205	Gln	Lys	Lys	Phe	Lys 210	723
	Asn	Ser	Thr	TAC	Ser 215	Arg	Ser	Ser	Val	Asp 220	Val	Leu	Туг	Thr	Phe 225	768 813
25	Ala	Asn	Cys	TCA Ser	Gly 230	Leu	Asp	Leu	lle	Phe 235	Gly	Leu	Asn	Ala	240	858
30	Leu	Arg	Thr	Ala	Asp 245	Leu	Glri	7rp	Asn	Ser 250	Ser	Asn	Ala	Gln	255	903
	Leu	cac	ASP	TYF	Cys 260 CCT	Ser	Ser	Lys	Gly	Tyr 265	ASD	Ile	Ser	Trp.	Glu 270 TYC	948
35	l eu	GLY	Asn	Glu	Pro 275 CAG	Asn.	Ser.	Phe GAA	Leu	Lys 280 TAT	Lys	Ala	Asp	Ile	Phe 285 AAA	993
40 ·	Ile	ASD	Gly	Ser	Gln 290 TCC	Leu	Gly	Glu	ASP	Tyr 295 GCA	Ile	Gln	TAT	His	Lys 300 CCT	1038
	GAT	GTI	GGT	Lys	305 CCT	CGA	AGA	AAG	ACG	310 GCT	AAG	ATG	CTG	AAG	315 AGC	1083
45	TTC	CTG	AAG	GIN GCT Ala	320 GGT	GGA	GAA	GTG	ATT	325 GAT	TCA	GTT	ACA	TGG	CAT	1128
	CAC	TAC	TAT	TIG Leu	335 AAT	GGA	CGG	ACT	GCT	340 ACC	AGG	GAA	GAT	TTT	345 CTA	1173
50	AAC	133	GAT	GTA Val	350 116	GAC	ATT	111	ATT	355 TCA	161	GTG	CAA	AAA	360 GTT	1218
55	TTC	CAG	GTG	GTT	365 GAG	AGC	ACC	AGG	CCT	370 GGC	AAG	AAG	GTC	TGG	375 TTA	1263
	Phe	Gln	Val	Val	380	ser	100	AFG	PFO	385	LYS	LYS	vat	irp	390	

				GCA Ala									1308
5				GGC									1353
10				ATA									1398
				CAT His									1443
15				TCT Ser									1488
				AGC Ser									1533
20				TGC Cys									1578
25				CTG Leu									1623
			Arg	CCC Pro									1668
30				CCT Pro									1713
35			Leu	GGT Gly		Leu					Asp		1758
			Pro	ATG Het		Pro					Ser		1803
40			Рго	TTC Phe		Ser							1848
	Ala	Lys	Ala	TGC Cys	TGA	AAA	TAA	AAT	ATA	CTA	GTC	CTG	1893
45	ACA	CTG											1899

## (2) INFORMATION FOR SEQ ID NO:16:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 594

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(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16

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5	ATTACTATAG GGCACGCGTG GTCGACGGCC CGGGCTGGTA TTGTCTTAAT GAGAAGTTGA 60 TAAAGAATIT TGGGTGGTTG ATCTCTTTCC AGCTGCAGTT TAGCGTATGC TGAGGCCAGA 120 FTTTTTTCAGG CAAAAGTAAA ATACCTGAGA AACTGCCTGG CCACAAGGACA ATCAGATTTT 180 GGCTGGCTCA AGTGACAAGC AAGTGTTAAT AAGCTAGATG GGAGAGGAAG GGATGAATAC 240 TCCATTGGAG GCTTTACTCG AGGGTCAGAG GGATACCCGG CGCCATCAGA ATGGGATCTG 300 GGAGTCGGAA ACCTGGGTT CCCACGAGAG CGCGCAGAAC ACCTGCGTCA GGAAGCCTGG 360 TCCGGGGAGGC CCGCGCGTGC TCCCCCGCG CTCCTCCCCG GGCGCTCCTC CCCAGGCCTC 420 CCGGGCGCGTT GGATCCCGGC CATCTCCGCA CCCTTCAAGT GGGTTGTGGGT GATTTCGTAA 480 GTGAACGTGA CCGCCACCGG GGGGAAAGCG ACCAAGGAAG TAGGAGAGAG CCGGGCAGGC 540 GGGGCGGGGT TGGATTGGGA GCAGTGGGA GAGAGGAAG GAGGAGTGGG AGGC 594
10	(2) INFORMATION FOR SEQ ID NO:17:
	(2) INI ONMATION FOR SEQ ID NO. 17.
	(i) SEQUENCE CHARACTERISTICS:
15	(A) LENGTH: 21 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17
	CCCCAGGAGC AGCAGCATCA G 21
25	(2) INFORMATION FOR SEQ ID NO:18:
	(i) SEQUENCE CHARACTERISTICS:
30	(A) LENGTH: 21 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18
35	
	AGGCTTCGAG CGCAGCAGCA 1 21
	(2) INFORMATION FOR SEQ ID NO:19:
40	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 22
	(B) TYPE: nucleic acid
45	(C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19
50	
	GTAATACGAC TCACTATAGG GC 22
	(2) INFORMATION FOR SEQ 10 NO:20:
55	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 19 (B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20 ACTATAGGGC ACGCGTGGT 19 (2) INFORMATION FOR SEQ ID NO:21: . 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 (B) TYPE: nucleic acid (C) STRANDEDNESS: single 15 (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21 20 . CTTGGGCTCA CCTGGCTGCT C 21 (2) INFORMATION FOR SEQ ID N0:22: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 (B) TYPE: nucleic acid (C) STRANDEDNESS: single 30 (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID ND:22 AGCTCTGTAG ATGTGCTATA CAC 23 35 (2) INFORMATION FOR SEQ ID NO:23: (i) SEQUENCE CHARACTERISTICS: 40 (A) LENGTH: 22 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 45 (xi) SEQUENCE DESCRIPTION: SEO ID NO:23 GCATCTTAGC CGTCTTTCTT CG 22 50 Claims 1. An isolated polynucleotide fragment comprising a polynucleotide sequence encoding a polypeptide having hepara-55

- nase catalytic activity, wherein said polypeptide shares at least 70% homology with SEQ ID NOs: 10 or 14 or a functional fragment thereof having heparanase catalytic activity.
- 2. The polynucleotide fragment of claim 1, wherein said polynucleotide sequence includes nucleotides 63 1691 of

SEQ ID NO: 9 or nucleotides 139 - 1869 of SEQ ID NO: 13.

- 3. The polynucleotide fragment according to claim 1 or 2, wherein said polynucleotide sequence includes nucleotides 63 721 of SEQ ID NO- 9.
- 4. The polynucleotide, fragment of claim 1, wherein said polynucleotide is as set forth in SEQ ID NO: 9 or 13.
- 5. The polynucleotide fragment of claim 1, wherein said polypeptide includes an amino acid sequence as set forth in SEQ ID NOs: 10 or 14.
- 6. A polynucleotide fragment comprising a polynucleotide sequence at least 70% homologous with SEQ ID NOs: 9 or 13, said polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.
- The polynucleotide fragment of any preceding claim, wherein said polynucleotide sequence is selected from the group consisting of double stranded DNA, single stranded DNA and RNA.
  - 8. A vector comprising a polynucleotide fragment according to any preceding claim.
  - 9. A host cell comprising an exogenous polynucleotide fragment according to any of claims 1 to 7.
  - 10. A recombinant protein, which is a polypeptide of 543 amino acids as set forth in SEQ ID NO: 10 with a calculated molecular weight of 61,192 daltons or a functional part thereof
- 11. A polypeptide of 592 amino acids as set forth in SEQ ID NO: 14 with a calculated molecular weight of 66,407 daltons or a functional part thereof.
  - 12. A pharmaceutical composition comprising, as an active ingredient, the isolated protein/polypeptide according to claim 10 or 11 and a pharmaceutically acceptable carrier.
- 13. A medical device containing, as an active ingredient, an isolated protein/potypeptide according to claim 10 or 11.
  - 14. A heparanase overexpression system comprising a cell overexpressing heparanase sharing at least 70% homology with SEQ ID Nos: 10 or 14 and encoded by a polynucleotide fragment as claimed in any one of claims 1 to 7.
- 15. A method of identifying a chromosome region harbouring a heparanase gene in a chromosome spread comprising the steps of:
  - (a) hybridizing the chromosome spread with a tagged polynucleotide probe according to claims 6 or 7,
  - (b) washing the chromosome spread, thereby removing excess of non-hybridized probe; and
  - (c) searching for signals associated with said hybridized tagged probe, wherein detected signals being indicative of a chromosome region harbouring a heparanase gene.

#### Patentansprüche

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- 1. Isoliertes Polynucleotidfragment, umfassend eine Polynucleotidsequenz, die ein Polypeptid codiert, das Heparanase-katalytische Aktivität besitzt, wobei das Polypeptid mindestens 70% Homologie mit SEQ ID NR:10 oder 14 oder einem funktionalen Fragment davon mit Heparanase-katalytischer Aktivität aufweist.
- Polynucleotidfragment nach Anspruch 1, wobei die Polynucleotidsequenz Nucleotide 63-1691 von SEQ ID NR:9 oder Nucleotide 139-1869 von SEQ ID NR:13 umfasst.
  - Polynucleotidfragment nach Anspruch 1 oder 2, wobei die Polynucleotidsequenz Nucleotide 63-721 von SEQ ID NR:9 umfasst.
  - 4. Polynucleotidfragment nach Anspruch 1, wobei das Polynucleotid wie in SEQ ID NR:9 oder 13 dargestellt ist.
  - 5. Polynucleotidfragment nach Anspruch 1, wobei das Polypeptid eine wie in SEQ ID NR:10 oder 14 dargestellte

Aminosäuresequenz umfasst.

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- 6. Polynucleotidfragment, umfassend eine Polynucleotidsequenz mit mindestens 70% Homologie mit SEQ ID NR:9 oder 13, wobei die Polynucleotidsequenz ein Polypeptid mit Heparanase-katalytischer Aktivität codiert.
- 7. Polynucleotidfragment nach einem der vorangegangenen Ansprüche, wobei die Polynucleotidsequenz ausgewählt ist aus einer Gruppe bestehend aus doppelsträngiger DNA, einzelsträngiger DNA und RNA.
- 8. Vektor, umfassend ein Polynucleotidfragment nach einem der vorangegangenen Ansprüche.
- 9. Wirtszelle, umfassend ein exogenes Polynucleotidfragment nach einem der Ansprüche 1 bis 7.
- 10. Rekombinantes Protein, welches ein Polypeptid mit 543 Aminosäuren ist, wie dargestellt in SEQ ID NR:10, mit einem berechneten Molekulargewicht von 61.192 Dalton, oder ein funktionelles Fragment davon.
- 11. Polypeptid mit 592 Aminosäuren, wie dargestellt in SEQ ID NR:14, mit einem berechneten Molekulargewicht von 66.407 Dalton, oder ein funktionelles Fragment davon.
- Arzneimittel, umfassend, als einen aktiven Bestandteil, das isolierte Protein/Polypeptid nach Anspruch 10 oder 11
   und einen pharmazeutisch verträglichen Träger.
  - Medizinprodukt, enthaltend, als einen aktiven Bestandteil, ein isoliertes Protein/Polypeptid nach Anspruch 10 oder 11.
- 25 14. Heparanase-Überexpressionssystem, umfassend eine Zelle, welche eine Heparanase überexprimiert, die mindestens 70% Homologie mit SEQ ID NR:10 oder 14 aufweist und durch ein Polynucleotidfragment gemäß einem der Ansprüche 1 bis 7 codiert wird.
- 15. Verfahren zur Identifizierung einer Chromosomenregion, die ein Heparanasegen enthält, in einer Chromosomen-30 ansammlung, umfassend die Schritte:
  - (a) Hybridisieren der Chromosomenansammlung mit einer markierten Polynucleotidsonde nach einem der Ansprüche 6 oder 7;
  - (b) Waschen der Chromosomenansammlung, wobei ein Überschuss von nicht hybridisierter Sonde entfernt wird; und
  - (c) Suchen nach Signalen, die mit der hybridisierten markierten Sonde assoziiert sind, wobei der Nachweis von Signalen auf eine Chromosomenregion hinweist, die ein Heparanasegen enthält.

### 40 Revendications

- Fragment polynucléotidique isolé comprenant une séquence polynucléotidique codant un polypeptide ayant une activité catalytique héparanase, dans lequel ledit polypeptide possède au moins 70% d'homologie avec SEQ ID NO: 10 ou 14 ou un fragment fonctionnel de celles-ci ayant une activité catalytique héparanase.
- 2. Fragment polynucléotidique selon la revendication 1, dans lequel la séquence polynucléotidique inclut les nucléotides 63 1691 de SEQ ID NO : 9 ou les nucléotides 139 -1869 de SEQ ID NO 13.
- Fragment polynucléotidique selon la revendication 1 ou 2, dans lequel la séquence polynucléotidique inclut les nucléotides 63 - 721 de SEQ ID NO : 9.
  - Fragment polynucléotidique selon la revendication 1, dans lequel ledit polynucléotide est tel que représenté dans SEQ ID NO: 9 ou 13.
- 55 5. Fragment polynucléotidique selon la revendication 1, dans lequel ledit polypeptide inclut une séquence d'acides aminés telle que représentée dans SEQ ID NO : 10 ou 14.
  - 6. Fragment polynucléotidique comprenant une séquence polynucléotidique au moins 70% homologue à SEQ ID

NOs: 9 ou 13, ladite séquence polynucléotidique codant un polypeptide ayant une activité catalytique héparanase.

- 7. Fragment polynucléotidique selon l'une quelconque des revendications précédentes, dans lequel ladite séquence polynucléotidique est choisie dans le groupe constitué d'ADN double-brin, d'ADN simple-brin et d'ARN.
- 8. Vecteur comprenant un fragment polynucléotidique selon l'une quelconque des revendications précédentes.
- 9. Cellule hôte comprenant un fragment polynucléotidique exogène selon l'une des revendications 1 à 7.

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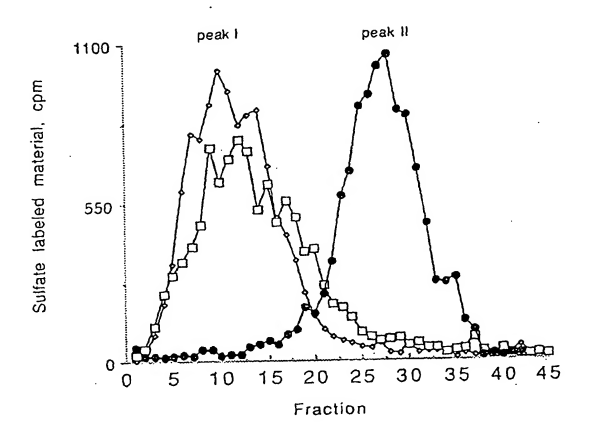
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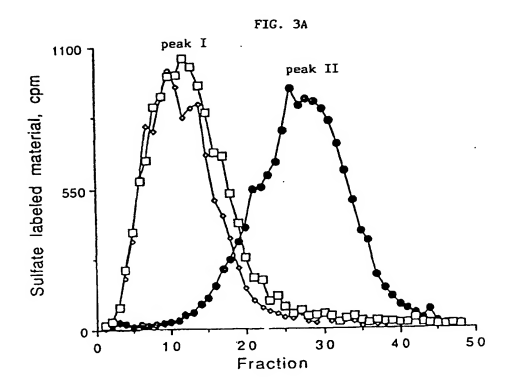
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- 10. Protéine recombinante, qui est un polypeptide de 543 acides aminés tel que représenté dans SEQ ID NO : 10 ayant un poids moléculaire calculé de 61 192 daltons, ou une partie fonctionnelle de celui-ci.
  - 11. Polypeptide de 592 acides aminés tel que représenté dans SEQ ID NO : 14 ayant un poids moléculaire calculé de 66 407 daltons, ou une partie fonctionnelle de celui-ci.
  - 12. Composition pharmaceutique comprenant, en tant qu'ingrédient actif, la(le) protéine/polypeptide isolé(e) selon la revendication 10 ou 11 et un support acceptable sur le plan pharmaceutique.
- 13. Dispositif médical contenant, en tant qu'ingrédient actif, un(e) protéine/polypeptide isolé(e) selon la revendication 20 10 ou 11.
  - 14. Système pour la sur-expression d'une héparanase, comprenant une cellule sur-exprimant une héparanase ayant au moins 70% d'homologie avec SEQ ID NO : 10 ou 14 et codée par un fragment polynucléotidique tel que revendiqué dans l'une des revendications 1 à 7.
  - 15. Méthode pour identifier une région chromosomique portant un gène héparanase à partir d'un étalement de chromosomes, comprenant les étapes consistant à :
    - (a) hybrider l'étalement de chromosomes avec une sonde polynucléotidique étiquetée selon les revendications 6 ou 7
      - (b) laver l'étalement de chromosomes, éliminant ainsi l'excès de sonde non-hybridée; et
    - (c) rechercher des signaux associés à ladite sonde étiquetée, les signaux détectés étant indicatifs d'une région chromosomique portant un gène héparanase.

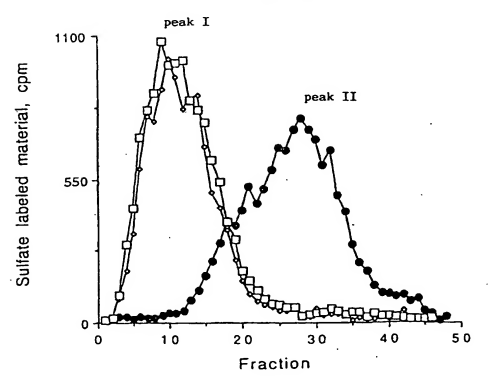
1	CTAGAGCT	TTCGA	CTCTCC	GC TC	SCGC	:GGC	AGC	TGC	CGG	GGG	<b>GA</b> (	SCA(	SCC.	AGG'	TGA	محو	CA
61	AGATGCTS M L																
121	CGCTG3GT L G																
181	ACCTGGAC L D																
241	CCATTSAC I D	eccaa A n	CCTGGC L A	CACO	GAC D	P CCG	CGG R	FTTC	CTC L	ATC I	CTC L	CTC L	3331 G	FTC: 5	rcc P	aaa K	3C L
301	TTCGTACC R T																
361	ACTTCCTAL F L															3C, Q	
421	CTCRASTC: Q V																
481	TACGGTTG																
541	TCAAGAACI K N	RGCRC: S T	CTACTO Y S	AAGI R	NAGC S	TCT S	GTA V	CAI D	GTG V	CT) L	Y Y	AC1	TTT F	GC) A	A N	273 <u>2</u>	c: S
601	CAGGACTS: G L	GACTT: D L	rapott 1 f	TGGC G	CTA L	aat N	GCG A	TTA L	1772 L	ag) R	laca T	IGC: A	GAS D	TTO L	FCA Q	GIG W	FR N
661	ACASTICIA S S I	RATGC' N A	TCASTT Q L	GCTC L	CTG L	GAC D	TAC	TGC C	TCT S	TCC S	iaac K	ecc G	TA? Y	iaac N	ia? I	7.TC	TT W
721	GGGRRADER E L G	BECAR B N	E 2	N	AGT S	TTC F	CTI L	aag K	AAG K-	GC1 A	GA? D	AT? I	e E	i I	n N	7,55 G	37 5
781	CGCAGTTAC		AGATTA D Y														
841	ATGUARAN A K I	CICTA! L Y	(F) COSTCC G P	TGAI D	GTT V	GGT G	CAG Q	E E	CG# R	AGZ R	K K	acd T	JCT A	R K	M M	3C7 L	K.
901	ASABOTEC: S F 1	CTGAA: L K	GCTGG A G	TGGA G	GAA E	GTG V	ATT I	GAT D	TCA S	GTT V	ACA T	TG: W	EA?	CAC S	TA Y	Y Y	TT L
291	TGRATUSAC N G																
1021	TTTCATCTC S S V																
1061	GGTTAFFA: L G	Garaci E t	AAGCTC S S	TGCA A	Y Y	GGA G	G G	G G	GCG A	P P	TTG L	icti L	S.	GA(	T T	CIT F	TG A
1141	CAGCTGGC: A G																
1201	TGATTAGG R R (	erroti V	KTTCTT F F	TGGP G	IGCA A	GGA G	AAC N	TAC Y	E E	TT; L	V V	GA7 D	gaj E	ra: N	er E	D D	TC 2
1261	CTTTACCTC	SATTAI D Y	TGGCT W L	ATCT S	CTT L	CTG L	TTC F	AAG K	aaj K	lt Të	576 V.	GGC G	ACC T	744 K	GT V	GIT L	AA M
1321	TGGCRAGCO A 5 V	TGCAI	AGGTTC G S	AAAG K	AGA R	agg R	AAG K	CTI L	CGA R	GT2 V	Y	CT?	CAT H	rte( C	CAC.	aaa N	CA T
1381	CTGACAATO D N i	CCAAGO PR	TATAA Y K	AGAA E	GGA G	GAT D	TTA L	ACT T	CTG	TA7 Y	GCC A	AT;	iaa: N	CTC L	CA E	TAR N	cs v
1441	TCACCAAS!	TACTTO L	CGGTT R L	ACCC P	TAT Y	CCT P	TT1 F	TCT S	rac N	AAC K	CA! Q	V V	GA?	R R	ATA Y	CCT L	TC L
1501	TAAGACCT1	ttggg; L g	ACTTCA P B	TGG? G	L L	CTT L	TCC S	XAA K	TC1 S	GTC V	CAP Q	CTC L	AAT N	ree: G	TCT L	AAC T	TC L
1561	TARREATES	etsga1 V D	rgatca D Q	AACC T	TTG L	CCA P	CCT P	TTA L	ATG M	GAJ E	R R	P P	CTC L	EG:	55C. P	as:s	2.A 5
1621	GTTCACTS:	FSCTTC	ECASC PA	TTTC F	TCA S	TAT Y	AGT S	TTT	TTI	'679 V	ATA I	iac:	AA: N	:GZ(	iaa K	:::: V	TG A

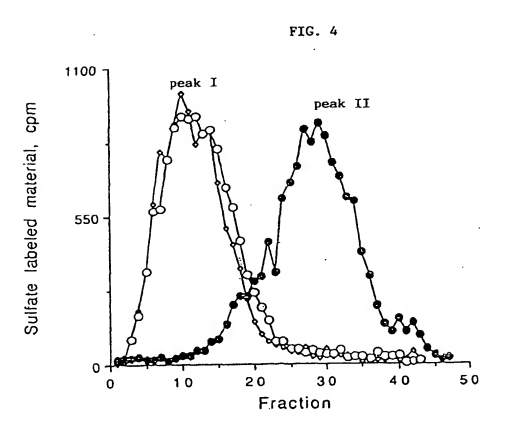
FIG. 2

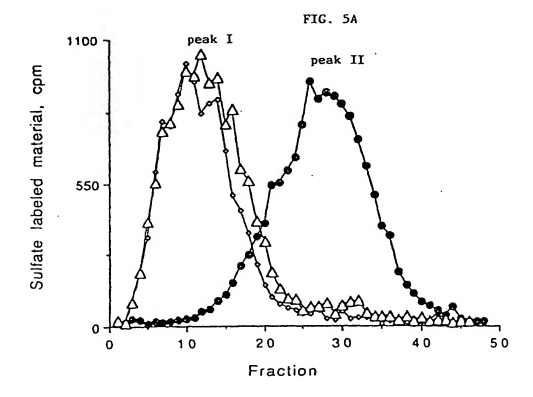




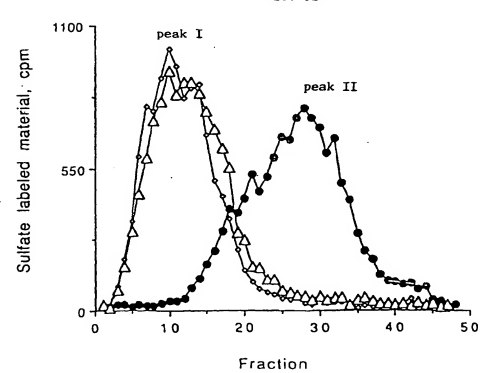


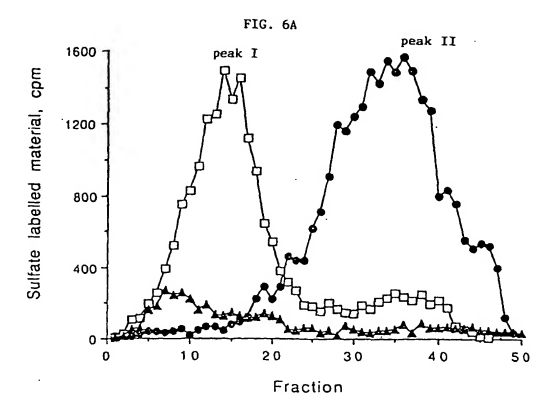












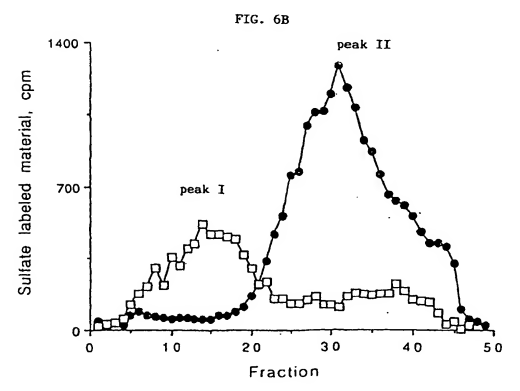


FIG. 7A

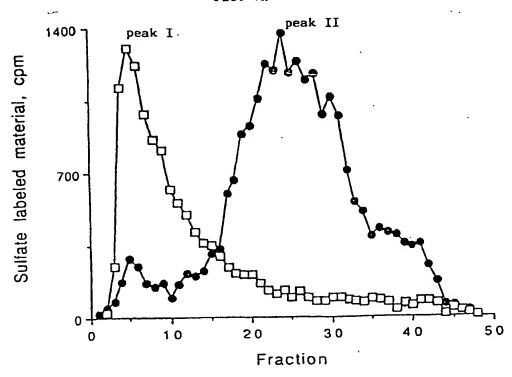
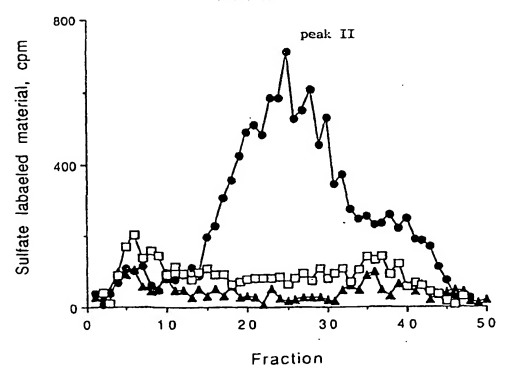
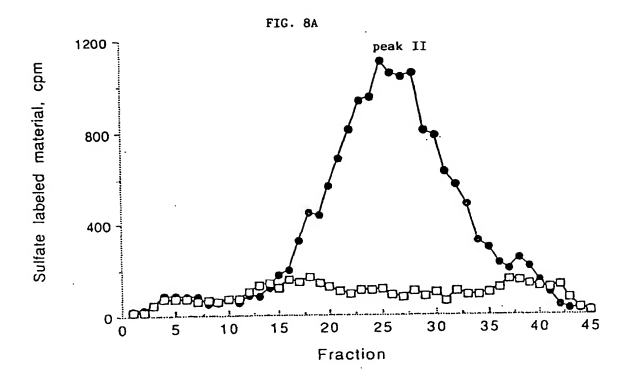


FIG. 7B





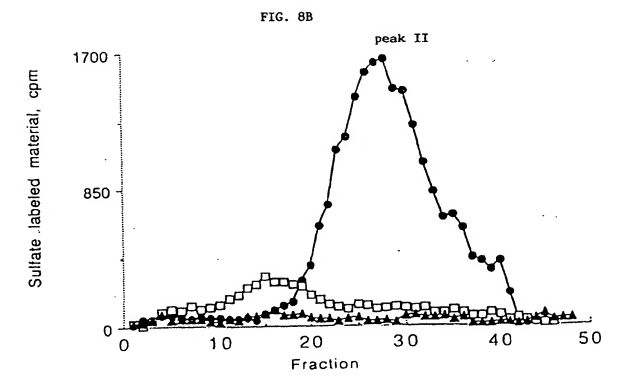


FIG. 9A

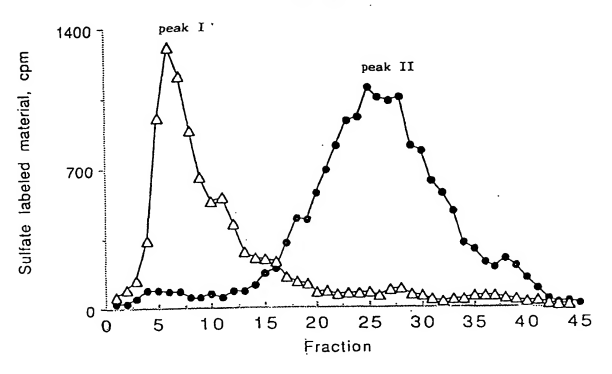
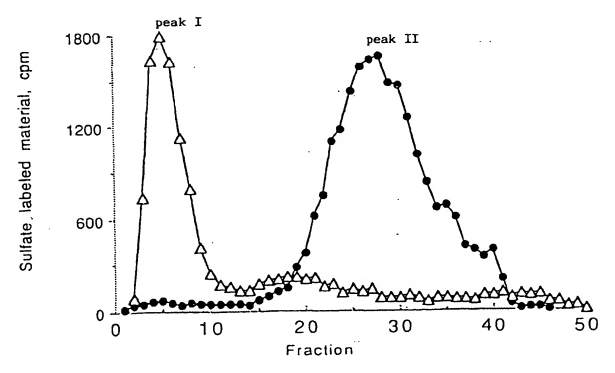
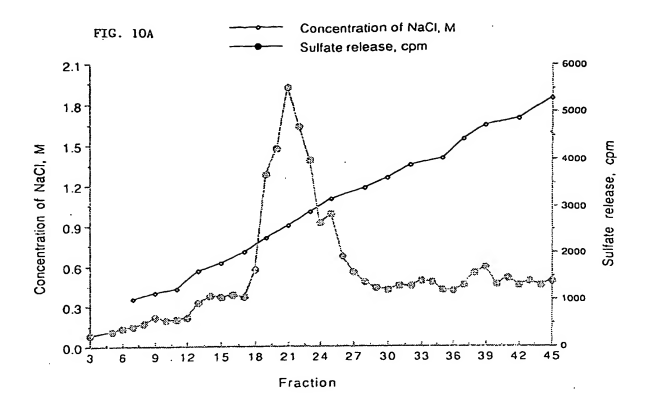
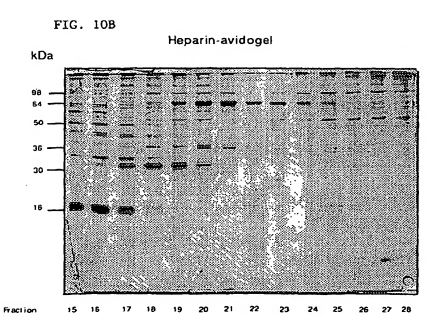
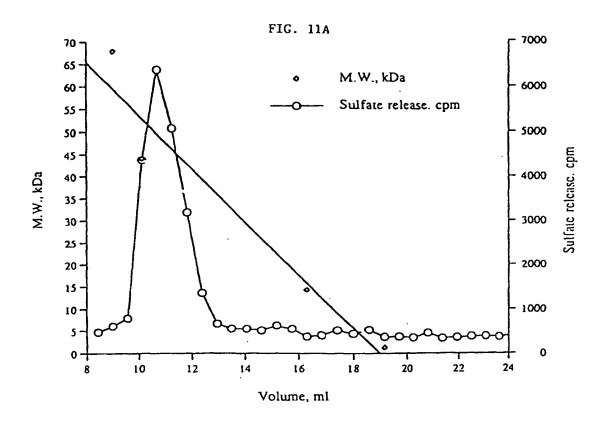


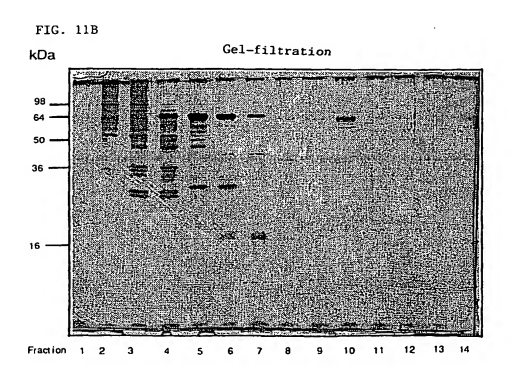
FIG. 9B

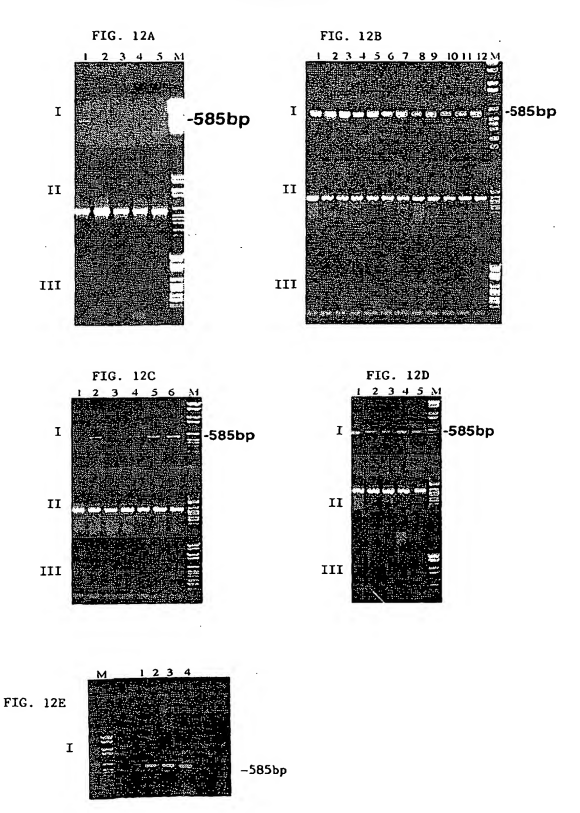












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# Fig. 13

mouse	CTGGCAAGAAGGTCTGGTTGGGAGAGACGAGCTCAGCTTACGGTGGCGGT	
human	CTGGCAAGAAGGTCTGGTTAGGAGAAACAAGCTCTGCATATGGAGGCGGA	
mouse	GCACCCTTGCTGTCCAACACCTTTGCAGCTGGCTTTATGTGGCTGGATAA	100
human	GCGCCCTTGCTATCCGACACCTTTGCAGCTGGCTTTATGTGGCTGGATAA	1165
mouse	ATTGGGCCTGTCAGCCCAGATGGGCATAGAAGTCGTGATGAGGCAGGTGT	150
human	ATTGGGCCTGTCAGCCCGAATGGGAATAGAAGTGGTGATGAGGCAAGTAT	1215
mouse	TCTTCGGAGCAGCAACTACCACTTAGTGGATGAAAACTTTGAGCCTTTA	200
human	TCTTTGGAGCAGGAAACTACCATTTAGTGGATGAAAACTTCGATCCTTTA	1265
mouse	CCTGATTACTGGCTCTCTCTTCTGTTCAAGAAACTGGTAGGTCCCAGGGT	250
human		1315
mouse	GTTACTGTCAAGAGTGAAAGGCCCAGACAGGAGCAAACTCCGAGTGTATC	300
human		1365
mouse	TCCACTGCACTAACGTCTATCACCCACGATATCAGGAAGGA	350
human	TTCATTGCACAACACTGACAATCCAAGGTATAAAGAAGGAGATTTAACT	1415
mouse	CTGTATGTCCTGAACCTCCATAATGTCACCAAGCACTTGAAGGTACCGCC	400
human		1465
mouse	TCCGTTGTTCAGGAAACCAGTGGATACGTACCTTCTGAAGCCTTCGGGGC	450
human		1515
mouse	CGGATGGATTACTTTCCAAATCTGTCCAACTGAACGGTCAAATTCTGAAG	500
human		1565
mouse	ATGGTGGATGAGCAGACCCTGCCAGCTTTGACAGAAAAACGTCTCCCCGC	550
human		1615
mouse	AGGAAGTGCACTAAGCCTGCCTGCCTTTTCCTATGGTTTTTTTT	600
human	AGGAAGTTCACTGGGCTTGCCAGCTTTCTCATATAGTTTTTTTT	1665
mouse	GAAATGCCAAAATCGCTGCTTGTATA <u>TGA</u> AAATAAAA 637	
human		

